World Health Organization Classification of Tumours

Pathology & Genetics

Head and Neck Tumours

Edited by Leon Barnes, John W. Eveson, Peter Reichart, David Sidransky

Diagnostic criteria, pathological features, and associated genetic alterations are described in a strictly disease-oriented manner. Sections on all WHO-recognized neoplasms and their variants include new ICD-O codes, incidence, age and sex distribution, location, clinical signs and symptoms, pathology, genetics, and predictive factors.

The book, prepared by 130 authors from 28 countries, contains 890 colour photographs, X-rays, computed tomography (CT), magnetic resonance (MR) images, charts, and more than 2900 references.
World Health Organization Classification of Tumours

International Agency for Research on Cancer (IARC)

Pathology and Genetics of Head and Neck Tumours

Edited by

Leon Barnes
John W. Eveson
Peter Reichart
David Sidransky

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International Academy of Pathology (IAP)

and the

Department of Pathology, University Hospital, Zurich, Switzerland


Members of the Working Group are indicated in the List of Contributors on page 371.
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CHAPTER 1

Tumours of the Nasal Cavity and Paranasal Sinuses

Although the nasal cavity and paranasal sinuses occupy a relatively small anatomical space, they are the site of origin of some of the more complex, histologically diverse group of tumours in the entire human body. These include neoplasms derived from mucosal epithelium, seromucinous glands, soft tissues, bone, cartilage, neural/neuroectodermal tissue, haematolymphoid cells and the odontogenic apparatus. Many of the tumours are similar to those found elsewhere in the body but a few, such as the olfactory neuroblastoma, are unique to this site.
**WHO histological classification of tumours of the nasal cavity and paranasal sinuses**

<table>
<thead>
<tr>
<th>Malignant epithelial tumours</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Squamous cell carcinoma</td>
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</tr>
<tr>
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<td>Leiomyma</td>
</tr>
<tr>
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</tr>
<tr>
<td>Basaloid squamous cell carcinoma</td>
<td>Schwanoma</td>
</tr>
<tr>
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<td>Neurofibroma</td>
</tr>
<tr>
<td>Adenosquamous carcinoma</td>
<td>Meningioma</td>
</tr>
<tr>
<td>Acantholytic squamous cell carcinoma</td>
<td></td>
</tr>
<tr>
<td>Lymphoepithelial carcinoma</td>
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</tr>
<tr>
<td>Sinonasal undifferentiated carcinoma</td>
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<table>
<thead>
<tr>
<th>Adenocarcinoma</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Intestinal-type adenocarcinoma</td>
<td>Chondrosarcoma</td>
</tr>
<tr>
<td>Non-intestinal-type adenocarcinoma</td>
<td>Mesenchymal chondrosarcoma</td>
</tr>
<tr>
<td></td>
<td>Osteosarcoma</td>
</tr>
<tr>
<td>Salivary gland-type carcinomas</td>
<td>Chordoma</td>
</tr>
<tr>
<td>Adenoid cystic carcinoma</td>
<td>Giant cell lesion</td>
</tr>
<tr>
<td>Acinic cell carcinoma</td>
<td>Giant cell tumour</td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td>Chordroma</td>
</tr>
<tr>
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</tr>
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<td>Clear cell carcinoma N.O.S.</td>
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</tr>
<tr>
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<td>Osteomyxoid fibroma</td>
</tr>
<tr>
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<td>Osteochondroma (exostosis)</td>
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<td>Polymorphous low-grade adenocarcinoma</td>
<td>Osteoid osteoma</td>
</tr>
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<td>Neuroendocrine tumours</td>
<td>Osteoblastoma</td>
</tr>
<tr>
<td>Typical carcinoid</td>
<td>Ameloblastoma</td>
</tr>
<tr>
<td>Atypical carcinoid</td>
<td>Nasal chondromesenchymal hamartoma</td>
</tr>
<tr>
<td>Small cell carcinoma, neuroendocrine type</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Benign epithelial tumours</th>
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</tr>
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<tbody>
<tr>
<td>Sinonasal papillomas</td>
<td></td>
</tr>
<tr>
<td>Inverted papilloma</td>
<td></td>
</tr>
<tr>
<td>(Schneiderian papilloma, inverted type)</td>
<td></td>
</tr>
<tr>
<td>Oncocytic papilloma</td>
<td></td>
</tr>
<tr>
<td>(Schneiderian papilloma, oncocytic type)</td>
<td></td>
</tr>
<tr>
<td>Exophytic papilloma</td>
<td></td>
</tr>
<tr>
<td>(Schneiderian papilloma, exophytic type)</td>
<td></td>
</tr>
<tr>
<td>Salivary gland-type adenomas</td>
<td></td>
</tr>
<tr>
<td>Pleomorphic adenoma</td>
<td></td>
</tr>
<tr>
<td>Myoepithelioma</td>
<td></td>
</tr>
<tr>
<td>Oncocytoma</td>
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<th>Soft tissue tumours</th>
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</thead>
<tbody>
<tr>
<td>Malignant tumours</td>
<td></td>
</tr>
<tr>
<td>Fibrosarcoma</td>
<td>8810/3</td>
</tr>
<tr>
<td>Malignant fibrous histiocytoma</td>
<td>8830/3</td>
</tr>
<tr>
<td>Leiomyosarcoma</td>
<td>8890/3</td>
</tr>
<tr>
<td>Rhabdomyosarcoma</td>
<td>8900/3</td>
</tr>
<tr>
<td>Angiosarcoma</td>
<td>9120/3</td>
</tr>
<tr>
<td>Malignant peripheral nerve sheath tumour</td>
<td>9540/3</td>
</tr>
<tr>
<td>Borderline and low malignant potential tumours</td>
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</tr>
<tr>
<td>Desmoid-type fibromatosis</td>
<td>8821/1</td>
</tr>
<tr>
<td>Inflammatory myofibroblastic tumour</td>
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<tr>
<td>Glomangiopericytoma</td>
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<tr>
<td>(Sinonasal-type haemangiopericytoma)</td>
<td>9150/1</td>
</tr>
<tr>
<td>Extramural solitary fibrous tumour</td>
<td>8815/1</td>
</tr>
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<table>
<thead>
<tr>
<th>Tumours of bone and cartilage</th>
<th>Malignant tumours</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Chondrosarcoma</td>
</tr>
<tr>
<td></td>
<td>Mesenchymal chondrosarcoma</td>
</tr>
<tr>
<td></td>
<td>Osteosarcoma</td>
</tr>
<tr>
<td></td>
<td>Chordoma</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>Benign tumours</th>
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<tbody>
<tr>
<td>Haematolymphoid tumours</td>
<td></td>
</tr>
<tr>
<td>Extramedullary plasmacytoma</td>
<td>9734/3</td>
</tr>
<tr>
<td>Extramedullary myeloid sarcoma</td>
<td>9930/3</td>
</tr>
<tr>
<td>Histiocytic sarcoma</td>
<td>9755/3</td>
</tr>
<tr>
<td>Langerhans cell histiocytosis</td>
<td>9751/1</td>
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<table>
<thead>
<tr>
<th>Neuroectodermal</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Primitive neuroectodermal tumour</td>
<td>9364/3</td>
</tr>
<tr>
<td>Olfactory neuroblasticoma</td>
<td>9522/3</td>
</tr>
<tr>
<td>Melanotic neuroectodermal tumour of infancy</td>
<td>9363/0</td>
</tr>
<tr>
<td>Mucosal malignant melanoma</td>
<td>8720/3</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Germ cell tumours</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Immature teratoma</td>
<td>9080/3</td>
</tr>
<tr>
<td>Teratoma with malignant transformation</td>
<td>9084/3</td>
</tr>
<tr>
<td>Sinonasal yolk sac tumour (endodermal sinus tumour)</td>
<td>9071/3</td>
</tr>
<tr>
<td>Sinonasal teratocarcinosarcoma</td>
<td></td>
</tr>
<tr>
<td>Mature teratoma</td>
<td>9080/0</td>
</tr>
<tr>
<td>Dermoid cyst</td>
<td>9084/0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Secondary tumours</th>
<th></th>
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<tbody>
<tr>
<td>Morphology code of the International Classification of Diseases for Oncology (ICD-O) (821) and the Systematized Nomenclature of Medicine (<a href="http://snomed.org">http://snomed.org</a>). Behaviour is coded /0 for benign tumours, /3 for malignant tumours, and /1 for borderline or uncertain behaviour.</td>
<td></td>
</tr>
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</table>
## TNM classification of carcinomas of the nasal cavity and paranasal sinuses

### T – Primary tumour

<table>
<thead>
<tr>
<th>T</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>TX</td>
<td>Primary tumour cannot be assessed</td>
</tr>
<tr>
<td>T0</td>
<td>No evidence of primary tumour</td>
</tr>
<tr>
<td>Tis</td>
<td>Carcinoma in situ</td>
</tr>
</tbody>
</table>

### Maxillary sinus

<table>
<thead>
<tr>
<th>T</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1</td>
<td>Tumour limited to the antral mucosa with no erosion or destruction of bone</td>
</tr>
<tr>
<td>T2</td>
<td>Tumour causing bone erosion or destruction, including extension into hard palate and/or middle nasal meatus, except extension to posterior antral wall of maxillary sinus and pterygoid plates</td>
</tr>
<tr>
<td>T3</td>
<td>Tumour invades any of the following: bone of posterior wall of maxillary sinus, subcutaneous tissues, floor or medial wall of orbit, pterygoid fossa, ethmoid sinuses</td>
</tr>
<tr>
<td>T4a</td>
<td>Tumour invades any of the following: anterior orbital contents, skin of cheek, pterygoid plates, infratemporal fossa, cribiform plate, sphenoid or frontal sinuses</td>
</tr>
<tr>
<td>T4b</td>
<td>Tumour invades any of the following: orbital apex, dura, brain, middle cranial fossa, cranial nerves other than maxillary division of trigeminal nerve V2, nasopharynx, clivus</td>
</tr>
</tbody>
</table>

### Nasal cavity and ethmoid sinus

<table>
<thead>
<tr>
<th>T</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>T1</td>
<td>Tumour restricted to one subsite of nasal cavity or ethmoid sinus, with or without bony invasion</td>
</tr>
<tr>
<td>T2</td>
<td>Tumour involves two subsites in a single site or extends to involve an adjacent site within the nasoethmoidal complex, with or without bony invasion</td>
</tr>
<tr>
<td>T3</td>
<td>Tumour extends to invade the medial wall or floor of the orbit, maxillary sinus, palate, or cribiform plate</td>
</tr>
<tr>
<td>T4a</td>
<td>Tumour invades any of the following: anterior orbital contents, skin of nose or cheek, minimal extension to anterior cranial fossa, pterygoid plates, sphenoid or frontal sinuses</td>
</tr>
<tr>
<td>T4b</td>
<td>Tumour invades any of the following: orbital apex, dura, brain, middle cranial fossa, cranial nerves other than V2, nasopharynx, clivus</td>
</tr>
</tbody>
</table>

### N – Regional lymph nodes

<table>
<thead>
<tr>
<th>N</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>NX</td>
<td>Regional lymph nodes cannot be assessed</td>
</tr>
<tr>
<td>N0</td>
<td>No regional lymph node metastasis</td>
</tr>
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### M – Distant metastasis

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<th>Description</th>
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<tr>
<td>MX</td>
<td>Distant metastasis cannot be assessed</td>
</tr>
<tr>
<td>M0</td>
<td>No distant metastasis</td>
</tr>
<tr>
<td>M1</td>
<td>Distant metastasis</td>
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### Stage grouping

<table>
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<th>Stage</th>
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<th>N</th>
<th>M</th>
</tr>
</thead>
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<td>0</td>
<td>Tis</td>
<td>N0</td>
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</tr>
<tr>
<td>I</td>
<td>T1</td>
<td>N0</td>
<td>M0</td>
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<td>II</td>
<td>T2</td>
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<td>III</td>
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<td>M0</td>
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<td>T3</td>
<td>N0, N1</td>
<td>M0</td>
</tr>
<tr>
<td>IVA</td>
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<td>M0</td>
</tr>
<tr>
<td></td>
<td>T4a</td>
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<td>Any N</td>
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<td>M0</td>
</tr>
<tr>
<td>IVC</td>
<td>Any T</td>
<td>Any N</td>
<td>M1</td>
</tr>
</tbody>
</table>

---

1. [947,2418].
3. The regional lymph nodes are the cervical nodes.
Anatomy

The nasal cavities are separated in the midline by the nasal septum. Each cavity is wide caudally, and narrow cranially. The roof of the nasal cavity is formed by the thin (0.5 mm) cribriform plate. The floor is the hard palate, formed by the palatine processes of the maxillae and the horizontal portions of the palatine bones. The lateral nasal wall contains the maxillary and ethmoid ostia, plus three or four turbinates. These turbinates are delicate scroll-like projections of bone and vascular soft tissue that become smaller as they ascend in the nasal cavity. They attach to the lateral nasal wall anteriorly, and have a free edge posteriorly. The turbinates are covered with a thick mucous membrane and contain a dense, thick-walled venous plexus. The upper margins of the nasal fossa are bound laterally by the superior nasal turbinate and adjacent lateral nasal wall, and medially by the nasal septum. This region is the olfactory recess and it has a yellowish epithelium, the olfactory mucosa (OM). This mucosa contains bipolar olfactory nerve fibers that cross through the cribiform plate. The terminal axons of the olfactory nerves extend to the free surface of the epithelium, where they expand into knob-like protrusions bearing cilia (olfactory cilia). Bowman’s glands, or olfactory glands, within the lamina propria appear similar to serous minor salivary glands.

The nasal cavity and paranasal sinuses are lined by Schneiderian mucosa, consisting of pseudostratified columnar ciliated epithelium with interspersed goblet cells. The lamina propria within the paranasal sinuses, especially the maxillary antrum, is loose and well vascularized, with seromucinous glands, and can easily become polypoid as a result of edema. The goblet cell component of the mucosal surface and seromucinous glands is variable. In chronic sinusitis, goblet cell hyperplasia can result in a papillary mucosal lesion. The major portion of the nasal septum is formed by the perpendicular plate of the ethmoid bone posteriorly and the septal cartilage anteriorly. The vomer completes the posterosuperior portion of the septum. The septum is lined by relatively thin, ciliated respiratory mucosa, which may regularly undergo squamous metaplasia. The underlying thin lamina propria, although containing seromucinous glands, is tethered to the septal cartilage, restricting reactive polyp formation.

The frontal sinus

These paired sinuses reside between the internal and external cranial tables and drain either via a nasofrontal duct into the anterior recess or more directly into the anterior infundibulum, or less often, into the anterior ethmoid cells, which in turn will open into the infundibulum of the bulla ethmoidalis.

Ethmoid complex

This paired complex of sinuses contains 3-18 cells that are grouped as anterior, middle, or posterior, according to the location of their ostia. There is an inverse relationship between the number and size of the cells. Generally, the posterior cells are both larger and fewer than the anterior cells. Each ethmoid labyrinth lies between the orbit and the upper nasal fossa. The left and right groups of ethmoid cells are connected in the midline by the cribiform plate (nasal roof) of the ethmoid bone. The cribiform plate is an important landmark in evaluation of sinonasal tumour stage - violation of the cribiform plate signifies direct extension of the tumour into the anterior cranial fossa. The crista galli is a distinctive pointed bony landmark that extends from the midline of the cribiform plate upward into the floor of the anterior cranial fossa. The perpendicular plate of the ethmoid bone extends downward from the cribiform plate to contribute to the nasal septum. The medial wall of each ethmoid labyrinth is formed by a thin lamella of bone from which arise the middle, superior, and supreme turbinates. The lateral ethmoid wall is formed by the thin lamina papyracea, which separates the ethmoid cells from the orbit. This is yet another important landmark for tumour staging. Tumour violation of the lamina papyracea may necessitate including the orbit and globe with the surgical resection. This area should be sampled in a maxillectomy specimen a) if the globe has not been removed (as the lamina papyracea represents the lateral orbital margin), or b) if orbital exenteration has been performed. The roof of the ethmoid complex is formed by a medial extension of the orbital plate of the frontal bone, which projects to articulate with the cribiform plate. This is often referred to as the fovea ethmoidalis.

Sphenoid sinus

The average adult sphenoid sinus measures 20 mm high, 23 mm long, and 17 mm wide. The relationship of the posterior extension of the sphenoid in relation to the sella turcica is variable. The sphenoid sinus septum is usually in the midline, and anteriorly aligned with the nasal septum. However, it can also deviate far to one side creating two unequal sinus cavities. With the exception of the sinus roof, the other sinus walls are of variable thickness depending on the degree of pneumatization. The sphenoid roof is thin, often measuring only 1 mm. (planum sphenoidale), and is vulnerable to perforation during surgery. The sinus roof relates to the floor of the anterior cranial fossa, anteriorly; the optic chiasm and
the sella turcica, posteriorly. The lateral sphenoid wall is related to the orbital apex, the optic canal, the optic nerve, and the cavernous sinus, containing the internal carotid artery. The sinus floor is the roof of the nasopharynx, and the anterior sinus wall is the back of the nasal fossa.

Maxillary sinus
The maxillary sinus lies within the body of the maxillary bone. Behind the orbital rims, each sinus roof/orbital floor slants obliquely upward so that the highest point of the sinus is in the posteromedial portion, lying directly beneath the orbital apex. The medial antral wall is the inferior or lateral wall of the nasal cavity (“party wall”). The curved posterolateral wall separates the sinus from the infratemporal fossa. The anterior sinus wall is the facial surface of the maxilla that is perforated by the infraorbital foramen below the orbital rim. The floor of the sinus is lowest near the second premolar and first molar teeth and usually lies 3-5 mm below the nasal floor. The lower expansion of the antrum is intimately related to dentition. The location of the maxillary sinus ostia, is high on the medial wall. They drain through the ethmoidal infundibulum and then the nasal fossa. This pattern of drainage in the erect position is accomplished by intact ciliary action. The maxillary hiatus is a bony window leading to the interior of the maxillary sinus. The hiatus is normally partially covered by portions of four bones: the perpendicular plate of the palatine bone, posteriorly; the lacrimal bone, anterosuperiorly; the inferior turbinate, inferiorly, and above the turbinate attachment, the uncinate process of the ethmoid bone.

Fig. 1.2 Malignancy of ethmoid and nasal cavity. Coronal CT with contrast. The tumour erodes the cribriform plate and fovea ethmoidalis (white arrowhead). The lamina papyracea is eroded (black arrowhead) but the fat (white arrow) medial to the medial rectus (MR) is normal. The orbital fat is not invaded. The margin (black arrow) of the tumour in the maxillary sinus is separable from the obstructed secretions because of different densities.

Fig. 1.3 Malignancy of upper nasal cavity invading orbital fat and extending intracranially. The tumour (white arrow) extends through the roof of the ethmoid and along the roof of the orbit (white arrowhead). The tumour bulges (black arrow) the periorbita near the medial rectus but breaks through into the orbital fat more superiorly (black arrowhead). Note that the tumour enhances intermittently and less intensely than the mucosa.

Etiology
Occupational exposure to wood dust, in particular to dust of hard woods such as beech and oak, is the main known risk factor for sinonasal cancer. The increase in risk (in the order of 5-50 fold) is strongest for adenocarcinomas and for cancers originating from the sinuses. The effect is present after 40 or more years since first exposure and persists after cessation of exposure. An increased risk of sinonasal cancer has been shown among workers in nickel refining and chromate pigment manufacture, but not among workers exposed to these metals in other processes, such as plating and welding. Among other suspected occupational carcinogens are formaldehyde, diisopropyl sulfide and dichloroethyl sulfide.

A relatively weak (relative risks in the range 2-5) but consistent association has been shown between tobacco smoking and sinonasal cancer, in particular squamous cell carcinoma. Exposure to Thorotrast, a radioactive contrast agent, represents an additional risk factor.

Imaging
Modern imaging plays a key role in the evaluation of sinonasal tumours [2423]. The anatomy of the lesion can be defined with the exact margins clearly delineated in almost every case. Imaging is a dominant factor in determining surgical approach and is an integral part of radiation therapy planning. Computed tomography (CT) and magnetic resonance imaging (MRI) provide significant information about the texture, the margins, the effect on bone and even the vascularity. In addition, some findings are typical for a particular diagnosis, and although biopsy is still required for ascertaining the nature of the lesion, the imaging appearance may help limit the list of differential diagnoses.

Staging and surgical planning
The spread of a sinonasal tumour intracranially or into the orbit and the relationship of tumour to the optic nerve and carotid artery are important features that can be delineated with imaging. Tumour can invade the orbit through the lamina papyracea or the roof of the maxillary sinus. Even if the bony wall is apparently destroyed, orbital fat may not be invaded [515]. A smooth bowing of the soft tissue interface with the orbital fat

Epidemiology
Carcinomas of the nasal cavity and paranasal sinuses account for 0.2-0.8% of all malignant neoplasms and 3% of those occurring in the head and neck [169,2378]. Sixty percent of sinonasal tumours originate in the maxillary sinus, 20-30% in the nasal cavity, 10-15% in the ethmoid sinus, and 1% in the sphenoid and frontal sinuses [2378]. Malignant neoplasms of this region may lead to significant morbidity and disfigurement.

The incidence of cancer of the nasal cavity and paranasal sinuses (sinonasal cancer) is low in most populations (<1.5/100,000 in men and <1.0/100,000 in women). Higher rates are recorded in Japan and certain parts of China and India. Squamous cell carcinomas are the commonest. Time trends have shown in most populations a stable incidence or a small decline in recent decades.

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suggests that the lesion is contained by periorbital fascia. Infiltration or irregularity of this margin suggests extension into the fat or true orbital invasion. The thin line of fat between the medial rectus muscle and the lamina papyracea is a key landmark in the evaluation of orbital extension of ethmoid neoplasms. The key landmarks for the assessment of intracranial extension of tumour are the roof of the ethmoid, the cribriform plate and the crista galli. Elevation or frank invasion of the dura may be evaluated using MRI.

A tumour in the maxillary sinus region may extend posteriorly and laterally through the bony wall into the pterygopalatine fossa and the infratemporal fossa. Tumour can invade the pterygopalatine fossa area either by direct extension or by following the nerves. From there, perineural extension of tumour in the foramen rotundum and Vidian canal may result in intracranial spread (516).

Tumour may spread from the sphenoid sinus region, laterally into the cavernous sinus through the very thin layer of bone separating these two structures. If the bone is intact, the tumour is likely contained within the sinus.

**Radiographic signs**

Bone changes can give an indication of the aggressiveness of a tumour (2038). In general, slowly growing lesions, such as Schneiderian papillomas, appear to push bone as they slowly remodel the osseous structure. More aggressive lesions, such as squamous cell carcinoma, can aggressively destroy bony walls leaving only a few remaining fragments (2425). Occasionally, however, malignant lesions can cause bowing rather than infiltrating destruction of bone (2424).

The integrity of the thin plates of bone in the ethmoid sinus as well as the bony walls of the sphenoid sinus and the bony nasal septum also suggests that malignancy is unlikely.

Mineralization can be seen in several tumours, such as ring-like calcifications in cartilage lesions as well as calcifications in olfactory neuroblastomas (2130). Meningioma can cause hyperostosis and can also calcify.

Tumour location plays a significant role in differential diagnosis. Tumours in the region of the cribriform plate and upper nasal cavity suggest diagnoses such as olfactory neuroblastoma or meningioma. Inverted Schneiderian papilloma occurs predominantly along the lateral wall of the nasal cavity or the medial maxillary sinus (534). In the lower maxilla, odontogenic lesions should be considered. Such lesions arise in the bone of the alveolar process and as they grow elevate the floor of the maxillary sinus. Fibro-osseous lesions enter the differential diagnosis when a radiodense lesion arises from or follows the contour of bone. Correlation of imaging studies with histologic appearance is crucial in the evaluation of bony lesions.

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**Fig. 1.4** Sinonasal polyps with multiple obstructed sinuses. **A** Axial T1 WI after intravenous gadolinium. The enhancing mucosa (arrowheads) lines the obstructed sphenoid sinus (S). The sinus was also dark on T2WI indicating high protein and long standing obstruction. **B** Coronal T1WI shows cascading polyps filling the nasal cavity. The secretions in the maxillary sinus (S) are dark and the lining mucosa (white arrowhead) is visible. The nasal septum (black arrow) is intact. Hard palate - (black arrowhead), minor salivary glands at roof of mouth (G), olfactory bulb (white arrow).

**Fig. 1.5** Sinonasal polyps. Coronal T2WI. The secretions in the sinus (S) are dark indicating high protein desiccated secretions. Compare to high signal of the edematous mucosa. Intact nasal septum (black arrow), crista galli (white arrow).
Squamous cell carcinoma

Definition
A malignant epithelial neoplasm originating from the mucosal epithelium of the nasal cavities or paranasal sinuses that includes a keratinizing and a non-keratinizing type.

ICD-O codes
Squamous cell carcinoma 8070/3
Verrucous carcinoma 8051/3
Papillary squamous cell carcinoma 8052/3
Basaloid squamous cell carcinoma 8083/3
Spindle cell carcinoma 8074/3
Adenosquamous carcinoma 8560/3
Acantholytic squamous cell carcinoma 8075/3

Synonyms
Keratinizing squamous cell carcinoma: squamous cell carcinoma.
Nonkeratinizing carcinoma: Schneiderian carcinoma, cylindrical cell carcinoma, transitional (cell) carcinoma, Ringertz carcinoma, respiratory epithelial carcinoma.

Epidemiology
Sinonasal squamous cell carcinoma is rare, accounting for <1% of malignant tumours and only about 3% of malignancies of the head and neck [169,2758]. The disease appears to be more common in Japan than in the West [2205]. It is extremely rare in children, and men are more commonly affected (about 1.5 times) than women. Patients are generally about 55-65 years of age [502,2758].

Etiology
Reported risk factors have included exposure to nickel, chlorophenols, and textile dust, prior Thorotrast instillation, smoking, and a history or concurrence of sinonasal (Schneiderian) papilloma. Human papillomavirus (HPV) has been found in some cases, especially those associated with inverted Schneiderian papilloma [303], but a definite etiologic role has not been clearly established. Formaldehyde, despite the results of animal experiments, has not been found to be a definite risk factor in humans [502,1443,1571,2205,2904].

Localization
Sinonasal squamous cell carcinomas occur most frequently in the maxillary sinus (about 60-70%), followed by the nasal cavity (about 12-25%), ethmoid sinus (about 10-15%) and the sphenoid and frontal sinuses (about 1%) [131,502]. Squamous cell carcinoma of the nasal vestibule should be considered a carcinoma of the skin rather than sinonasal mucosal epithelium [2566].

Clinical features
Symptoms include nasal fullness, stuffiness, or obstruction; epistaxis; rhinorrhea; pain; paraesthesia; fullness or swelling of the nose or cheek or a palatal bulge; a persistent or non-healing nasal sore or ulcer; nasal mass; or, in advanced cases, proptosis, diplopia, or lacrimation [131,502,2758]. Radiologic studies such as CT scan or MRI may delineate the extent of the lesion, the presence of bony invasion, and extension to neighbouring structures such as the orbit, pterygopalatine or infratemporal spaces.

Macroscopy
Sinonasal squamous cell carcinomas may be exophytic, fungating, or papillary; friable, haemorrhagic, partially necrotic, or indurated; demarcated or infiltrative.

Tumour spread and staging
Nasal cavity carcinomas can spread to adjacent sites in the nasal cavity or to the ethmoid sinus, or can extend to involve

Fig. 1.6 A Non-keratinizing papillary squamous cell carcinoma. Multiple complex papillary projections lined by thickened epithelium. Lymphocytic response is present at the pushing border of infiltration. B Squamous cell carcinoma, non-keratinizing. Islands of cohesive tumour cells invading into the underlying stroma. Surface carcinoma in-situ is seen.
the contralateral nasal cavity, bone, maxillary sinus, palate, skin and soft tissues of the nose, lip, or cheek, cribriform plate, or cranial cavity. Maxillary sinus carcinomas may spread to the nasal cavities, palate, other paranasal sinuses, skin or soft tissues of the nose or cheek, orbit, cranial contents, or the pterygopalatine and infratemporal spaces (131,2418). Lymph node metastases are less common than in squamous cell carcinomas of other sites in the head and neck.

**Histopathology**

**Keratinizing squamous cell carcinoma**
This tumour is histologically identical to squamous cell carcinomas of other mucosal sites in the head and neck. There is histologic evidence of squamous differentiation, in the form of extracellular keratin or intracellular keratin (pink cytoplasm, dyskeratotic cells) and/or intercellular bridges. Tumour cells are generally apposed to one another in a “mosaic tile” arrangement. The tumour may be arranged in nests, masses, or as small groups of cells or individual cells. Invasion occurs as blunt projections or ragged, irregular strands. There is often a desmoplastic stromal reaction. The carcinomas may be well, moderately, or poorly differentiated.

**Non-keratinizing (cylindrical cell, transitional) carcinoma**
This is a distinctive tumour of the sinonasal tract characterized by a plexiform or ribbon-like growth pattern. It invades into the underlying tissue with a smooth, generally well-delineated border. Therefore, definite evidence of stromal invasion may be difficult to appreciate, although a degree of invasion by irregular small nests or strands may be present. There is typically a lack of maturation in the epithelial nests or ribbons, as in transitional cell carcinoma of the urinary tract, which this tumour subtype resembles. Cytologic atypia is present to a significant degree. As its name implies, this tumour does not generally evince histologic evidence of keratinization, although some degree may be seen. When keratinization is significant, there is morphologic overlap with keratinizing squamous cell carcinoma. Occasional mucus-containing cells can be seen. The tumour may be moderately or poorly differentiated; the latter type is difficult to recognize as squamous, and must be differentiated from olfactory neuroblastomas or neuroendocrine carcinomas.

**Variants of squamous cell carcinoma**
Variants of squamous cell carcinoma are rare in the sinonasal tract. They are similar to the analogous tumours occurring with greater frequency in other sites in the head and neck and are more completely described in the corresponding sections. **Verrucous carcinoma** of the nasal and paranasal sinuses is very rare. It is a low-grade variant of squamous cell carcinoma characterized by a papillary or warty exophytic mass of very well-differentiated, keratinized epithelium (899,1955, 2118,2278). The maxillary sinus is the most common site, followed by the nasal fossa. Rare nasopharyngeal lesions have encroached on the nasal sinus (1199,1872).

**Papillary squamous cell carcinoma** (2488) is an exophytic squamous cell carcinoma with a papillary configuration composed of thin fingers of tumour surrounding fibrovascular cores. **Basaloid squamous cell carcinoma** is uncommon in the sinonasal tract (2786). It is an aggressive variant of squamous cell carcinoma that is characterized by rounded nests of cytologically highly atypical and mitotically-active basaloid epithelial cells, with high nuclear/cytoplasmic ratios and hyperchromatic nuclei. There is often comedo-type necrosis. A pseudoglandular or strand-like arrangement, reminiscent of the architecture of an adenoid cystic carcinoma, is often present, as is the production of basement membrane-like material. Squamous differentiation is invariably present, either in basaloid nests, as separate foci of tumour, or as surface epithelial carcinoma or carcinoma in-situ. **Spindle cell carcinoma** is characterized by a biphasic pattern of squamous cell carcinoma as well as a generally much larger component of malignant spindled cells, reminiscent of a sarcoma. The squamous component may be scant or even inapparent on light microscopy. In the latter circumstance, immunohisto-
chemical or ultrastructural evidence of epithelial differentiation is required for the diagnosis. The spindle cell component is characteristically immunohistochemically vimentin-positive, and keratin positivity may be scant, difficult to demonstrate, or even absent.

Adenosquamous carcinoma is uncommon in the sinonasal tract, and is more completely described in the sections on oral and laryngeal tumours. Briefly, it is generally considered as a variant of squamous cell carcinoma in which a surface mucosal component of squamous cell carcinoma is present. There is also a component of carcinoma with definite glandular differentiation in the form of ductules or tubules, often intimately admixed with the squamous cell carcinoma. The mere presence of intracellular mucin is not sufficient for the diagnosis.

Acantholytic squamous cell carcinoma is exceedingly rare in the sinonasal tract.

Precursor lesions
Precursor lesions for sinonasal squamous cell carcinomas are considerably less well defined than for oral or laryngeal carcinomas. The sinonasal Schneiderian (inverted) papilloma appears to be a precursor lesion; the frequency of association has been estimated at about 10% [173]. Although squamous metaplasia may precede the development of sinonasal squamous carcinoma, a predisposing role for such metaplasia in the development of carcinoma has not been clearly established.

Prognosis and predictive factors
Patients with nasal squamous cell carcinomas generally present earlier than patients with maxillary cancers and, not surprisingly, fare better than the latter group. Nasal squamous cell carcinomas rarely metastasize to lymph nodes, and recurrences, when they occur, do so quickly [131]. Advanced local disease worsens the prognosis. The overall 5-year survival for nasal squamous cell carcinomas is about 60%. Squamous carcinomas of the maxillary sinus have a more ominous prognosis. They are likely to be large and extensive when diagnosed. Prognosis correlates with stage. Patients with the non-keratinizing type of carcinoma tend to do better than those with the keratinizing type [502]. The overall 5-year survival of patients with maxillary sinus squamous carcinoma is about 42% [131].

Fig. 1.8 Schneiderian papilloma with keratinization is associated with an area of malignant transformation into a squamous cell carcinoma with severe cytologic atypia.
**Lymphoepithelial carcinoma**

**Definition**
Lymphoepithelial carcinoma is a poorly differentiated squamous cell carcinoma or histologically undifferentiated carcinoma accompanied by a prominent reactive lymphoplasmacytic infiltrate, morphologically similar to nasopharyngeal carcinoma.

**ICD-O code** 8082/3

**Synonyms**
Undifferentiated carcinoma; undifferentiated carcinoma with lymphocytic stroma; undifferentiated carcinoma of nasopharyngeal type; lymphoepithelioma-like carcinoma

**Epidemiology**
Sinonasal lymphoepithelial carcinoma is rare, and most reported cases have originated from Southeast Asia, where nasopharyngeal carcinoma is also prevalent (1216,1480,1558,2910). It affects adults in the fifth to seventh decades, and there is a male predominance of approximately 3:1.

**Etiology**
Nearly all sinonasal lymphoepithelial carcinomas show a strong association with Epstein-Barr virus (EBV) (801,1216,1480,1558,2910).

**Localization**
Sinonasal lymphoepithelial carcinomas are more common in the nasal cavity than in the paranasal sinuses, although both sites may be involved simultaneously. The tumours may show local invasion of the palate, orbit, and base of skull.

**Clinical features**
Patients present with nasal obstruction, bloody nasal discharge or epistaxis. Intracranial extension of tumour may cause proptosis and cranial nerve palsy (1216,1480). There may be cervical lymph node and/or distant metastasis at presentation. Examination and biopsy of the nasopharynx is required to exclude loco-regional spread from a primary nasopharyngeal carcinoma.

**Histopathology**
The tumour infiltrates the mucosa in the form of irregular islands and sheets, usually without a desmoplastic stroma. The tumour cells possess relatively monotonous vesicular nuclei with prominent nucleoli. The cytoplasm is lightly eosinophilic, with indistinct cell borders, resulting in a syncytial appearance. The tumour cells may also appear plump spindly, with streaming of nuclei. Intraepithelial spread of tumour may sometimes be seen in the overlying epithelial lining. Necrosis and keratinization are usually not evident. The tumour is infiltrated by variable numbers of lymphocytes and plasma cells. In general, the inflammatory infiltrate is less prominent than that seen in nasopharyngeal carcinoma. In some cases, the inflammatory cells may even be sparse (1216,1480). The epithelial nature of the tumour can be confirmed by immunostaining for pan-cytokeratin and epithelial membrane antigen. EBV encoded RNA (EBER) is strongly expressed by the tumour cells in most cases (801,1216,1480,1558,2910).

**Differential diagnosis**
Sinonasal lymphoepithelial carcinoma must be distinguished from the vastly more aggressive sinonasal undifferentiated carcinoma (SNUC). The presence of lymphoplasmacytic infiltrates, although helpful, cannot be relied on solely in making the distinction. SNUC is characterized by tumour cells with nuclear pleomorphism, high mitotic rate and frequent necrosis. EBV status is also helpful since SNUC, except for rare cases from Asians, are EBV-negative (1216,1480,1558). Other important differential diagnoses are malignant melanoma and non-Hodgkin lymphoma.

**Prognosis and predictive factors**
The tumour responds favourably to local-regional radiotherapy even in the presence of cervical lymph node metastasis (623,1216,1480). Distant metastasis (most often to bone), however, is associated with a poor prognosis.

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**Fig. 1.9** Primary lymphoepithelial carcinoma of the nasal cavity. A The intimate intermingling of the carcinoma cells with lymphoid cells imparts a lymphoma-like appearance. B Large carcinoma cells with indistinct cell borders, vesicular nuclei and prominent nucleoli are admixed with numerous small lymphocytes.
Sinonasal undifferentiated carcinoma

Definition
A highly aggressive and clinicopathologically distinctive carcinoma of uncertain histogenesis that typically presents with locally extensive disease. It is composed of pleomorphic tumour cells with frequent necrosis, and should be differentiated from lymphoepithelial carcinoma and olfactory neuroblastoma.

ICD-O code 8020/3

Synonym
Anaplastic carcinoma

Epidemiology
The tumour is rare, with fewer than 100 reported cases. The age range is broad (third to ninth decade), and the median age is in the sixth decade [350,1216]. There is a male predominance (2-3:1).

Etiology
The neoplasm is typically negative for Epstein-Barr virus [350,1216]. Some cases have occurred after prior radiation therapy for nasopharyngeal carcinoma [1216].

Localization
The nasal cavity, maxillary antrum, and ethmoid sinus are typically involved alone or in combination. The neoplasm also commonly extends to other contiguous sites.

Clinical features
Patients have multiple nasal/paranasal sinus symptoms, usually of relatively short duration, including nasal obstruction, epistaxis, proptosis, periorbital swelling, diplopia, facial pain, and symptoms of cranial nerve involvement.

Macrosopy
The tumour is usually larger than 4 cm. It is fungating, with poorly defined margins, bone destruction, and invasion of adjacent structures [2038].

Tumour spread and staging
In addition to involvement of multiple sinuses, the neoplasm destroys sinus walls and orbital bones. Penetration into the cranial cavity is frequent. Less often, there is extension into the nasopharynx or oral cavity. The tumour can metastasize to cervical lymph nodes and distant sites (such as liver, lung, bone) [1216].

Histopathology
Sinonasal undifferentiated carcinoma forms nests, lobules, trabeculae and sheets, in the absence of squamous or glandular differentiation. Severe dysplasia of the overlying surface epithelium has been noted in a few instances. The nuclei are medium to large-sized, surrounded by small amounts of eosinophilic cytoplasm that lacks a syncytial quality. The nucleoli are variable in size, but most often, they are single and prominent. The mitotic rate is very high and there is often prominent tumour necrosis and apoptosis. Lymphovascular invasion is often prominent.

Immunohistochemistry
The carcinoma is immunoreactive for pan-cytokeratins and simple keratins (CK7, CK8 and CK19), but not CK4, CK5/CK6 and CK14 [801]. Less than half of the cases have been reported to be positive for epithelial membrane antigen, neuron specific enolase, or p53 [350]. The tumour is negative for CEA, while positivity for synaptophysin, chromogranin, or S100 protein is only rarely observed [350,1216].

Electron microscopy
Ultrastructurally, cells with occasional small desmosomes and rare dense core granules have been noted [819].

Histogenesis
This is a tumour of uncertain histogenesis, but with unique clinicopathologic characteristics. It should be differentiated from other specific types of carcinoma and non-epithelial tumours with round cells.

Prognosis and predictive factors
Despite aggressive management, the prognosis is poor, with median survival of less than 18 months [350,1216], and 5-year survival of less than 20% [856]. Recent results suggest that more promising outcome may be achieved by combining chemoradiation and radical resection [1802].
Adenocarcinoma

These are glandular malignancies of the sinonasal tract, excluding defined types of salivary gland carcinoma. Two main categories are recognized: (1) intestinal-type adenocarcinoma, and (2) non-intestinal-type adenocarcinoma, which can be further divided into low-grade and high-grade subtypes. Overall, adenocarcinomas and salivary-type carcinomas comprise 10-20% of all sinonasal primary malignant tumours.

**Intestinal-type adenocarcinomas**

**Definition**
A primary malignant glandular tumour of the nasal cavity and paranasal sinuses histologically resembling adenocarcinoma or adenoma of the intestines, or exceptionally normal small intestinal mucosa.

**ICD-O code**
8144/3

**Synonyms**
Colonic-type adenocarcinoma, enteric-type adenocarcinoma.

**Epidemiology**
The frequency of intestinal type adenocarcinomas (ITACs) among primary sinonasal malignancies is difficult to ascertain. Most series report a pronounced male predominance, possibly because of occupational exposure. Patients have ranged in age from 12 to 86 years at the time of diagnosis (mean 58 years) [124].

**Etiology**
The causal relationship of wood dust and leather dust with the development of sinonasal ITACs has been established by several epidemiological studies from different countries [1594]. In this setting, dust particle size is important because those smaller than 5 µm reach the lower respiratory tract, while larger particles are accumulated in the nasal mucosa. However, the carcinogens involved in the onset of ITACs in wood workers and leather workers have not yet been clearly identified. Biologically active substances which can be present in wood and leather dusts include alkaloids, saponins, stilbenes, aldehydes, quinones, flavonoids, resins, oil, steroids, terpenes, fungal proteins, and tannins [1341]. Association has also been reported for agricultural workers, food manufacturers, and motor-vehicle drivers among men, and for textile occupations among women [1443].

**Localization**
ITACs involve the ethmoid sinus, nasal cavities and maxillary sinus in approximately 40%, 27% and 20% of cases, respectively. In the nasal cavities, the inferior and middle turbinates are the sites of predilection. For larger destructive lesions it may be impossible to ascertain the exact site of origin. Advanced tumours tend to invade the orbit, the pterygopalatine and infratemporal fossae, and the cranial cavity. About 10% of cases show lymph node involvement at presentation [124,1341,2234].

**Clinical features**
Most patients present with unilateral nasal obstruction, rhinorrhea and epistaxis. Advanced tumours may cause pain, neurologic disturbances, exophthalmos and visual disturbances.

**Imaging**
Computed tomography (CT) and magnetic resonance imaging (MRI) are used for diagnosis of early lesions, defining the extent of disease and detection of early recurrence. CT best shows sites of bone destruction, while MRI best delineates soft tissue extension [1537].

**Fig. 1.11** Intestinal-type adenocarcinoma  A Well differentiated intestinal type adenocarcinoma shows a papillary growth pattern and occasional tubular glands.  B Higher power view of a moderately differentiated intestinal type adenocarcinoma, showing glandular structures formed by cylindrical and goblet cells.
Macroscopy
ITACs present as an irregular exophytic pink or white mass bulging in the nasal cavity or paranasal sinus, often with a necrotic friable appearance. Some lesions are gelatinous.

Histopathology
Two classifications of ITACs have been proposed. Barnes divided these tumours into 5 categories: papillary, colonic, solid, mucinous and mixed. Kleinsasser and Schroeder divided ITACs into four categories: papillary tubular cylinder cell (PTCC) types I-III (I = well-differentiated, II = moderately-differentiated, III = poorly-differentiated) {799,804,1333}, alveolar goblet type, signet-ring type and transitional type. Either classification is acceptable, but for simplicity the Barnes classification is preferred and will be the one utilized in this description. The most common histologic types seen in association with wood workers as well as in sporadic cases are the papillary and colonic types {124,1333}. The papillary type (papillary tubular cylinder cell I or well-differentiated adenocarcinoma), which accounts for approximately 18% of cases, shows a predominance of papillary architecture with occasional tubular glands, minimal cytologic atypia, and rare mitotic figures. The colonic type (papillary tubular cylinder II or moderately-differentiated adenocarcinoma), representing approximately 40% of cases, shows a predominance of tubulo-glandular architecture, rare papillae, increased nuclear pleomorphism and mitotic activity. The solid type (papillary tubular cylinder III or poorly-differentiated adenocarcinoma), representing approximately 20% of cases, shows a loss of differentiation, characterized by solid and trabecular growth with isolated tubule formation, marked increase in number of smaller cuboidal cells with nuclear pleomorphism, round vesicular nuclei, prominent nucleoli and increased mitotic figures. Analogous to colonic adenocarcinoma, some ITACs are predominantly comprised of abundant mucus and are classified as the mucinous type. The mucinous type (alveolar goblet cell and signet ring) includes two growth patterns. In one pattern, there are solid clusters of cells, individual glands, signet ring cells, short papillary fronds with or without fibrovascular cores; mucin is predominantly intracellular and a mucomyxoid matrix may be present. The other pattern shows the presence of large, well-formed glands distended by mucus and extracellular mucin pools {799,804,1333}. In the latter type, pools of extracellular mucin are separated by thin connective tissue septa creating an alveolar type pattern. Predominantly cuboidal or goblet tumour cells are present in single layers at the periphery of mucus lakes. Mucus extravasation may elicit an inflammatory response that can include multinucleated giant cells. {799}. The mixed type (transitional) is composed of an admixture of two or more of the previously defined patterns. Irrespective of the histologic type, ITACs histologically simulate normal intestinal mucosa and may include villi, Paneth cells, enterochromaffin cells and muscularis mucosae {1739}. In rare instances, the tumour is so well differentiated that it is composed of well-formed villi lined by columnar cells resembling normal resorptive cells; in some cases, bundles of smooth muscle cells resembling muscularis mucosae may also be identified under the villi.

Immunohistochemistry
ITACs are diffusely positive for epithelial markers including pancytokeratin, epithelial membrane antigen, B72.3, Ber-EP4, BRST-1, Leu-M1, and human milk fat globule (HMFG-2) {1687}. They show CK20 positivity (73%) and variable CK7.

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<td>Papillary-type</td>
</tr>
<tr>
<td>Colonic-type</td>
</tr>
<tr>
<td>Solid-type</td>
</tr>
<tr>
<td>Mucinous type</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Mixed</td>
</tr>
</tbody>
</table>

$^a$PTCC, papillary tubular cylinder cell
$^b$Survival data derived from Kleinsasser and Schroeder {1333}
reactivity (43% to 93% of cases) [800]. CDX-2, a nuclear transcription factor involved in the differentiation of intestinal epithelial cells and diffusely expressed in intestinal adenocarcinomas, is commonly expressed in ITACs [800]. Information on CEA staining in ITACs is conflicting [1687,2660]. Scattered or groups of chromogranin-positive cells are frequently identified [1687]; these neuroendocrine cells may express a variety of hormone peptides, including serotonin, cholecystokinin, gastrin, somatostatin and leu-enkephalin [163].

Electron microscopy
ITAC demonstrates features of the intestinal epithelium [163]. Columnar cells present regular microvilli with cores of microfilaments that combine to form a band that inserts into the zonula adherens of the junctional complexes. Glycocalyx bodies as characteristic of intestinal-type epithelium may be identified between the microvilli. Endocrine cells with neurosecretory granules, Paneth cells with large exocrine granules, and goblet cells containing several mucin droplets in the apical cytoplasm are present in variable numbers.

Precursor lesions
The frequent presence of squamous metaplasia and/or dysplasia of the sinonasal epithelium in the vicinity of the tumour impairs mucociliary clearance, resulting in prolonged contact of carcino- genic substances with the mucosa [2789].

Histogenesis
It has been hypothesized that ITAC derives from a stem cell capable of undergoing differentiation into various type of epithelial cells (resorptive cells, goblet cells, neuroendocrine cells, Paneth cells) [1739].

Genetics
Genetic data are limited [2012,2013, 2218,2829]. K-RAS or H-RAS mutation has been detected in only about 15% of cases [2012,2218]. TP53 mutations are reported in 18-44% of cases [2013,2829]; mutations consist more frequently of C:G to A:T transitions and involve the CpG dinucleotides. Other gene alterations include loss of heterozygosity (LOH) at 17p13 and 9q21, and promoter methylation of p14(ARF) and p16(INK4a). A close association between TP53, p14(ARF) and p16(INK4a) gene deregulation has been found in tumours from individuals occupationally exposed to dusts [2013].

Prognosis and predictive factors
Sinonasal ITACs are generally locally aggressive tumours with frequent local failure (about 50% of cases), whereas metastasis to cervical lymph nodes and spread to distant sites are infrequent (about 10% and 20%, respectively) [124,799,804,1333]. The 5-year cumulative survival rate is around 40%, with most deaths occurring within 3 years. Since most patients present with advanced local disease, clinical staging generally has no relevant prognostic significance.

The histologic subtype has been identified as indicative of clinical behaviour in different series [124,799,804,1333]. The papillary type (papillary tubular cylinder cell adenocarcinoma) has a more indolent course, with little tendency to distant spread (5-year survival rate of about 80%). Conversely, the solid type (papillary tubular cylinder cell adenocarcinoma) and mucinous type adenocarcinoma have a very poor survival. Other factors that have been associated with a more aggressive behaviour are: H-RAS mutation, chromogranin expression and c-erbB-2 expression [855,1687,2012].

Although it has been suggested that ITACs occurring in occupational exposed individuals have a better prognosis than sporadic ITACs [124], this has not been confirmed in other reports [799].

Sinonasal non-intestinal-type adenocarcinoma

ICD-O code 8140/3

Synonyms
Sinonasal low-grade adenocarcinoma, terminal tubulius adenocarcinoma, sinonasal tubulopapillary low-grade adenocarcinoma.

Definition
Adenocarcinomas arising in the sinonasal tract that are not of minor salivary gland origin and do not demonstrate histopathologic features of the sinonasal intestinal-type adenocarcinoma. These adenocarcinomas are divided into low- and high-grade subtypes.

Epidemiology
Sinonasal non-intestinal-type adenocarcinomas predominantly occur in adults but have been identified over a wide age range from 9-80 years [1044]. The average patient age at presentation of low-grade adenocarcinomas is 53 years while that of high-grade ones is 59 years [1044]. There is a slight male predominance for the low-grade adenocarcinomas but a more marked male predilection in the high-grade ones [1044].

Etiology
There are no known occupational or environmental etiological factors.

Localization
The low-grade non-intestinal-type adenocarcinomas predilect to the ethmoid sinus (to a lesser extent as compared with the intestinal-type), and the high-grade non-intestinal-type adenocarcinomas predilect to the maxillary sinus.

Fig. 1.13 Sinonasal high-grade non-intestinal-type adenocarcinoma. Solid areas of the tumour show marked nuclear pleomorphism as well as an area of comedo-type necrosis.
Either tumour type may also originate in the nasal cavity, other paranasal sinuses, or in multiple sinonasal sites in various combinations. 

**Clinical features**

For low-grade adenocarcinomas, patients primarily present with nasal obstruction and epistaxis. Pain is an infrequent feature. The duration of symptoms ranges from 2 months to five years, with a median of 5.5 months. For high-grade adenocarcinomas, the presenting symptoms include nasal obstruction, epistaxis, pain and facial deformity (e.g., proptosis). The duration of symptoms ranges from two weeks to five years with a median of 2.5 months.

**Macroscopy**

The appearance varies, including well demarcated to poorly-defined and invasive, flat to exophytic or papillary growths with a tan/white to pink colour and a friable to firm consistency.

**Histopathology**

The low-grade non-intestinal-type adenocarcinomas are circumscribed or invasive, and have a glandular or papillary growth. Numerous uniform small glands or acini are arranged in a back-to-back or coalescent pattern with little or no intervening stroma. Occasionally, large, irregular cystic spaces can be seen. The glands are lined by a single layer of non-ciliated columnar cells with uniform, round nuclei, granular eosinophilic cytoplasm. The cells vary from orderly linear arrangement to stratification with loss of nuclear polarity.

The high-grade non-intestinal-type adenocarcinomas are invasive tumours with a predominantly solid growth pattern, but glandular and papillary patterns can also be present. These tumours are characterized by moderate to marked cellular pleomorphism, high mitotic activity, including atypical forms, and necrosis.

**Prognosis and predictive factors**

The treatment for sinonasal non-intestinal-type adenocarcinomas is complete surgical excision generally via a lateral rhinotomy; depending on the extent and histology of the neoplasm, the surgery varies from local excision to more radical procedures (maxillectomy, ethmoidectomy and additional exenterations). Radiotherapy may be utilized for extensive disease or for higher-grade neoplasms. The low-grade neoplasms have an excellent prognosis, while high-grade neoplasms have a dismal prognosis with a 3-year survival rate of only approximately 20%.

![Fig. 1.14 Sinonasal (mucosal) non-intestinal-type adenocarcinoma. A Complex glandular growth and focal papillary architecture. B The glands are lined by a single layer of cuboidal to columnar appearing cells with uniform, round nuclei, single small identifiable nucleoli and eosinophilic appearing cytoplasm.](image)

![Fig. 1.15 Sinonasal low-grade non-intestinal-type adenocarcinoma. A Complex glandular growth including back-to-back glands lacking an intervening fibrovascular stroma is characteristically seen. B The glands are comprised of a single layer of nonciliated columnar cells with uniform, round nuclei, granular eosinophilic cytoplasm. The cells vary from orderly linear arrangement to stratification with loss of nuclear polarity.](image)
Tumours of the nasal cavity and paranasal sinuses

Salivary gland neoplasms of the sinonasal tract are uncommon, and the majority are malignant [1039]. For details see Chapter 5 on tumours of salivary glands.

**ICD-O codes**
- Adenoid cystic carcinoma 8200/3
- Acinic cell carcinoma 8550/3
- Mucoepidermoid carcinoma 8430/3
- Epithelial-myoepithelial carcinoma 8562/3
- Clear cell carcinoma 8310/3

**Adenoid cystic carcinoma**
Adenoid cystic carcinoma is the most frequent malignant salivary gland-type tumour of the sinonasal tract. The age range is from 11-92 years [1039]. The majority develop in the maxillary sinus (about 60%) and nasal cavity (about 25%) [130]. The disease is often insidious, and symptoms include nasal obstruction, epistaxis, and pain, paraesthesia or anaesthesia. Swelling of the palate or face, and loosening of the teeth may be the presenting symptom. Many tumours are large and extensively infiltrative at the time of diagnosis. These tumours can be difficult to detect on plain film radiographs and often extend widely through bone before there is radiographical evidence of osseous destruction. In addition, the true extent of tumour spread is often underestimated by imaging techniques. The long-term prognosis is poor and the 10-year survival rate is only 7% [2444]. Most patients die as a result of local spread rather than metastatic disease [2799].

**Acinic cell carcinoma**
Acinic cell carcinoma is rare in the sinonasal tract and cases have been reported in the nasal cavity [996,1950, 2014,2244,2698] and maxillary sinuses [829,2860]. The signs and symptoms are non-specific but they include nasal obstruction and epiphora.

**Mucoepidermoid carcinoma**
Mucoepidermoid carcinomas are rare at this site, and should be distinguished from the more aggressive variants of squamous cell carcinoma, especially adenosquamous carcinoma [1039, 1291,2588].

**Epithelial-myoepithelial carcinoma**
Epithelial-myoepithelial carcinoma is rare in the sinonasal tract. Cases have been reported to involve the nasal septum, nasal cavity and maxillary sinus [1011,1221,1450,2506]. Signs and symptoms are non-specific but have

---

**Fig. 1.16** Adenoid cystic carcinoma. **A** CT scan shows a nasal sinus mass focally extending into the bone. **B** The tumour (T) is inhomogenous and extends from the maxillary sinus into the infratemporal (black arrowhead) and the pterygopalatine fossa (black arrow). Perineural spread follows the Vidian cana (white arrowhead).

**Fig. 1.17** Adenoid cystic carcinoma of nasal cavity. **A** Solid as well as tubular growth patterns are seen. **B** An intact surface mucosa overlying the cribiform and cystic patterns of a sinonasal adenoid cystic carcinoma. **C** A high power illustrating the relatively bland nuclear appearance with an intermediate to high nuclear to cytoplasmic ratio. Palisading is noted, along with small gland or tubule formation, in addition to the larger cyst-like spaces.
included the formation of polypoid masses and nasal obstruction.

**Clear cell carcinoma**
Clear cell carcinoma, N.O.S., of the sinonasal tract is rare (1757,1874) and it is important to exclude metastatic renal clear cell carcinoma (1664,2918). Microscopically, these tumours consist of closely packed, polygonal clear cells arranged in sheets and theques. They contain glycogen but no mucin.

**Other tumours**
A variety of other salivary gland-type carcinomas have been rarely reported in the nasal cavity and paranasal sinuses. These include: malignant myoepithelioma (2918), carcinoma ex pleomorphic adenoma (435), polymorphous low-grade adenocarcinoma (1536) and basal cell adenocarcinoma (785).

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**Table 1.2 Sinonasal glandular tumours**

<table>
<thead>
<tr>
<th>Tumour types</th>
<th>No of cases</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pleomorphic adenoma</td>
<td>73</td>
<td>23%</td>
</tr>
<tr>
<td>Oncocytic tumours</td>
<td>7</td>
<td>2%</td>
</tr>
<tr>
<td>Low-grade adenocarcinoma (including acinic cell carcinoma)</td>
<td>67</td>
<td>21%</td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td>17</td>
<td>5%</td>
</tr>
<tr>
<td>Adenoid cystic carcinoma</td>
<td>54</td>
<td>17%</td>
</tr>
<tr>
<td>High grade adenocarcinoma</td>
<td>93</td>
<td>30%</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>311</strong></td>
<td><strong>100%</strong></td>
</tr>
</tbody>
</table>

* Modified from Hefnner (1039)
Neuroendocrine tumours are very rare in the nasal cavity, paranasal sinuses or nasopharynx. The recognizable types are typical carcinoid, atypical carcinoid and small cell carcinoma neuroendocrine type. It is unclear whether large cell neuroendocrine carcinoma that corresponds to the pulmonary counterpart occurs in these sites. There are also rare cases that do not fit these categories, and the diagnostic label “neuroendocrine carcinoma, not otherwise specified” may be applied.

**Carcinoid tumour**

**ICD-O codes**
- Typical carcinoid: 8240/3
- Atypical carcinoid: 8249/3

Typical and atypical carcinoids of the nasal cavity and paranasal sinuses are exceedingly rare, possibly because they are under-reported or have been included under other non-descriptive categories, such as “neuroendocrine carcinoma” [1676,2007,2384,2776]. They are otherwise similar to carcinoids in other sites.

Patients have ranged in age from 13-65 years, and present with nasal obstruction, epistaxis and/or facial pain. Most tumours arise in the nasal cavity but may extend into adjacent sinuses. A patient with two carcinoids - nasal and pulmonary - has been described [2384]. Another individual with the Multiple Endocrine Neoplasia Type I (MEN1) has been reported to have a carcinoid of the sphenoid sinus [2776].

Paucity of cases and lack of significant follow-up preclude definitive statements about the prognosis. The tumours are at least locally aggressive.

**Small cell carcinoma, neuroendocrine type (SCCNET)**

**Definition**
Small cell carcinoma, neuroendocrine type is a high-grade carcinoma composed of small to intermediate sized cells resembling those of small cell carcinoma of pulmonary or extrapulmonary origin. Necrosis, large numbers of apoptotic cells, high mitotic rate, and lack of neurofibrillary stroma are microscopic hallmarks of this tumour.

**ICD-O code**
8041/3

**Synonyms**
- Small cell carcinoma, small cell neuroendocrine carcinoma
- Oat cell carcinoma
- Poorly differentiated neuroendocrine carcinoma

**Epidemiology**
SCCNET of the sinonasal tract is a rare tumour with no sex, racial, or geographic predilection and no known association with smoking or radiation. The age range is from 26-77 years with a mean of 49 years.

**Localization**
SCCNET most commonly arise in the superior or posterior nasal cavity, and often extend into the maxillary or ethmoid sinuses. Primary tumours of the maxillary or ethmoid sinuses without nasal involvement can be seen in approximately 45% of cases. Secondary involvement of the nasopharynx is present in a minority of patients. Advanced tumours may invade the skull base, orbit, or brain.

**Clinical features**
The most common symptoms are epistaxis and nasal obstruction, followed by facial pain, palpable facial mass, and exophthalmos. Rare tumours have shown elevated serum levels of ACTH, calci-
Neuroendocrine carcinomas

Tonyin, pro-gastrin releasing peptide (pro-GRP), or antidiuretic hormone with syndrome of antidiuretic hormone and hyponatremia [1259,1901,2042].

Pathology
See Chapter 3 under “Neuroendocrine Neoplasms of the Larynx.” An important differential diagnosis is olfactory neuroblastoma.

Prognosis and predictive factors
SCCNET are aggressive tumours with a poor prognosis and frequent local recurrence and distant metastasis despite multimodal therapy. Among twenty reported patients [849,1259,1358,1901,2009,2042,2134,2153,2728,2742], twelve (60%) died of disease, three (15%) were alive with no evidence of disease, four were alive with disease (20%), and one died of other causes. In a study of extrapulmonary small cell carcinomas [845], which included seven cases involving the paranasal sinuses, the median survival of 14 patients with primary head and neck small cell carcinomas was only 14.5 months [845]. Follow-up data have shown a local recurrence rate of 45% and a distant metastasis rate of 35%. Common sites of metastases include cervical lymph nodes, lung, liver, bone marrow, and vertebrae.
Schneiderian papillomas

The ectodermally derived ciliated respiratory mucosa that lines the nasal cavity and paranasal sinuses, so-called Schneiderian membrane, gives rise to three morphologically distinct types of papillomas. These are referred to individually as inverted, oncocytic, and exophytic papillomas or, collectively, as Schneiderian papillomas.

As a group, the Schneiderian papillomas are uncommon, representing only 0.4-4.7% of all sinonasal tumours [1423].

Inverted papilloma (Schneiderian papilloma, inverted type)

Definition
A papilloma derived from the Schneiderian membrane in which the epithelium invaginates into and proliferates in the underlying stroma.

ICD-O code 8121/1

Synonyms
Inverting papilloma, Schneiderian papilloma, papillomatosis

Epidemiology
Inverted papillomas are two to five times more common in males, and are found primarily in the 40-70 year age group. They are distinctly uncommon in children.

Table 1.3 Distribution of Schneiderian papillomas

<table>
<thead>
<tr>
<th>References</th>
<th>Total cases</th>
<th>Inverted</th>
<th>Oncocytic</th>
<th>Exophytic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hyams [1158]</td>
<td>315</td>
<td>149</td>
<td>10</td>
<td>156</td>
</tr>
<tr>
<td>Michaels and Young [1714]</td>
<td>191</td>
<td>139</td>
<td>16</td>
<td>36</td>
</tr>
<tr>
<td>Buchwald et al. [302]</td>
<td>82</td>
<td>58</td>
<td>5</td>
<td>19</td>
</tr>
<tr>
<td>Sarkar et al. [2246]</td>
<td>35</td>
<td>24</td>
<td>9</td>
<td>2</td>
</tr>
<tr>
<td>Weiner et al. [2737]</td>
<td>105</td>
<td>82</td>
<td>2</td>
<td>21</td>
</tr>
<tr>
<td>Total</td>
<td>728 (100%)</td>
<td>452 (62%)</td>
<td>42 (6%)</td>
<td>234 (32%)</td>
</tr>
</tbody>
</table>

Fig. 1.27 Inverted papilloma. Coronal CT. The tumour bows the bone. The calcification (white arrowhead) may represent a sclerotic fragment of inferior turbinate.

Fig. 1.28 Inverted papilloma. A Specimen removed intact. Note the opaque yellow-tan nodular polypoid appearance. B Cut surface of the lesion shown in A. Close inspection shows well-demarcated islands of epithelium which extend endophytically into the stroma.

Etiology
Although a viral origin of inverted papillomas has long been suspected, viral inclusions have never been unequivocally demonstrated by light or electron microscopy. In addition, they are almost invariably negative when stained for human papillomavirus (HPV) by the immunoperoxidase technique. HPV genomes, however, have been demonstrated in inverted papillomas by in situ hybridization or the polymerase chain reaction, particularly HPV 6 and 11, sometimes HPV 16 and 18, and exceptionally, HPV 57. The frequency of finding the virus by these specialized techniques is highly variable, ranging anywhere from 0-100% [127]. In a collective review of 341 inverted papillomas evaluated for the presence of HPV by a variety of sophisticated molecular techniques, 131 (38%) were positive. Whether the virus is a passenger or etiologically related to the papilloma is unclear [1596]. Epstein-Barr virus (EBV) DNA has been identified in 65% of inverted papillomas by polymerase chain reaction (PCR), raising the possibility that this virus might be involved in its pathogenesis [1596]. A subsequent study utilizing in-situ hybridization found no evidence of EBV in the tumour cells, suggesting that the reported PCR positivity might be related to the presence of EBV-positive lymphocytes in the tissues [842]. There is no known association of inverted papilloma with allergy, inflammation, smoking, noxious environmental agents or occupation [1158].
Localization
Inverted papillomas characteristically arise from the lateral nasal wall in the region of the middle turbinate or ethmoid recesses, and often extend secondarily into the sinuses, especially the maxillary and ethmoid and, to a lesser extent, the sphenoid and frontal. Isolated lesions of the paranasal sinuses without nasal involvement however, do occur. Almost none arise primarily on the nasal septum {1297}. Exceptionally, inverted papillomas may arise in sites other than the sinonasal tract. They have been recorded in the middle ear-mastoid {2757}, pharynx {2499}, nasopharynx {81}, and lacrimal sac {2217}. It has been suggested that ectopic migration of the Schneiderian membrane during embryogenesis could account for these aberrant papillomas in sites contiguous with the sinonasal tract {1158}. Whether all of these ectopic cases are bona fide inverted papillomas is uncertain. Although overwhelmingly unilateral, rare cases of bilateral inverted papillomas have been described {211}. Such occurrence, however, should always arouse the suspicion of septal erosion and perforation from unilateral disease.

Imaging
Findings on imaging vary with the extent of disease. Early on, there may be only a soft tissue density within the nasal cavity and/or paranasal sinuses. Later, with more extensive disease, unilateral opacification and thickening of one or more of the sinuses is common, as are expansion and displacement of adjacent structures. Pressure erosion of bone may also be apparent and must be distinguished from the destructive invasion associated with malignancy, such as de novo carcinoma or carcinoma arising in and/or associated with an inverted papilloma.

Clinical features
Signs and symptoms
Nasal obstruction is the most common presenting symptom. Other manifestations include nasal drainage, epistaxis, anosmia, headaches (especially frontal), epiphora, proptosis, and diplopia. Pain, on the other hand, is an uncommon initial complaint, occurring in only about 10% of all cases. When present, it should always arouse suspicion of secondary infection or malignant change.

On physical examination, inverted papillomas present as pink, tan, or grey; non-translucent; soft to moderately firm, polypoid growths with a convoluted or wrinkled surface.

Imaging
Findings on imaging vary with the extent of disease. Early on, there may be only a soft tissue density within the nasal cavity and/or paranasal sinuses. Later, with more extensive disease, unilateral opacification and thickening of one or more of the sinuses is common, as are expansion and displacement of adjacent structures. Pressure erosion of bone may also be apparent and must be distinguished from the destructive invasion associated with malignancy, such as de novo carcinoma or carcinoma arising in and/or associated with an inverted papilloma.
Histopathology

Inverted papillomas are composed exclusively or almost exclusively of hyperplastic ribbons of basement membrane-enclosed epithelium that grow endophytically into the underlying stroma. Infrequently, a minor exophytic component may be seen. The epithelium is multilayered, usually 5-30 cells thick, and formed of squamous or ciliated columnar (respiratory epithelial) cells admixed with mucocytes. Nonkeratinizing squamous or transitional-type epithelium tends to predominate, and is frequently covered by a single layer of ciliated columnar cells. An occasional case may be composed almost entirely of respiratory epithelium. Gradations between these two extremes are not uncommon, resulting in a transitional epithelium reminiscent of that seen in the urinary tract. All of these epithelial types may be present in the same lesion, and their proportions may vary widely in different lesions or even in different areas of the same papilloma. Mitoses are not numerous and, if present at all, are seen primarily in the basal and parabasal epithelium. Ten to 20% of inverted papillomas may show focal surface keratinisation, and 5-10% varying degrees of dysplasia [127]. These are not necessarily signs of malignancy, but they should alert the pathologist of the need of thorough evaluation of the papilloma. The stroma ranges from dense and fibrous to loose and myxoid, with or without an inflammatory component. The inflammatory cells, especially neutrophils, often transmigrate through the epithelium. Basement membrane thickening is not typically seen. Normal-appearing seromucinous glands are sparse to absent, because the neoplastic epithelium uses the ducts and glands as scaffolds to extend into the stroma. As inverted papillomas enlarge, they may obstruct the drainage of nearby sinuses. As a result, it is not uncommon to also find ordinary nasal polyps in inverted papilloma specimens. They can usually be identified grossly by their more myxoid appearance and the fact that they will transmit, whereas inverted papilloma will not.

Rarely, an inverted papilloma will exhibit focal surface changes reminiscent of a verruca vulgaris; that is, it shows focal papillary squamous epithelial hyperplasia with marked keratosis and/or parakeratosis, with a prominent granular cell layer, and often contains numerous vacuolated cells suggestive of koilocytes. Although this might be a viral effect, immunohistochemical stains for HPV are invariably negative. When this change is observed, the diagnosis of “inverted papilloma with focal verrucous hyperplasia” is appropriate and the patient should be followed closely for possible development of carcinoma, either verrucous carcinoma or squamous cell carcinoma.

Inverted papilloma and carcinoma

Inverted papillomas are occasionally complicated by carcinomas, especially squamous cell carcinoma and, to a much lesser extent, verrucous, mucoepidermoid, spindle and clear cell carcinomas, as well as adenocarcinomas. The incidence of malignant change in individual series of inverted papillomas has ranged from 2-27% [127]. In a collective review of 1390 inverted papillomas reported in the literature, 150 (11%) were associated with carcinoma and, of these, 61% of the carcinomas were synchronous and 39% metachronous [127]. For metachronous carcinomas, the mean interval from onset of the inverted papilloma to the development of the carcinoma is 63 months (range, 6 months to 13 years) [1477]. Carcinomas complicating inverted papilloma vary from well to poorly differentiated and exhibit a broad range of behaviour. Some are in situ and of little consequence, whereas others are locally aggressive or may even metastasize. The carcinomas may actually arise within the papilloma, as evidenced by a gradation of histological changes ranging from dysplasia to carcinoma in-situ to frankly invasive carcinoma; whereas in others, the carcinoma is merely associated with a histologically bland inverted papilloma. Staining for CD44s may be helpful in identifying a malignant component. It is diffusely expressed in typical inverted papillomas, whereas its expression is reduced or absent in the associated carcinomatous component [1175].

There is no correlation between the number of local recurrences of an inverted papilloma and the subsequent development of carcinoma. There is some evidence, however, to suggest that HPV 16 and 18 may be more carcinogenic than HPV 6 and 11 [1334]. Preliminary data suggest that alterations in TP53, manifested by an increased protein expression or genetic mutation, can be used to predict which lesions are at risk for malignant change [715,765].

Differential diagnosis

The differential diagnoses include nasal polyp with squamous metaplasia, respiratory epithelial adenomatoid hamartoma (REAH) and invasive carcinoma. Nasal polyps with squamous metaplasia show thickening and hyalinization of the basement membrane, a prominent component of normal seromucinous glands and, often, a large number of stromal inflammatory cells. These features are typically absent in inverted papilloma. In addition, the surface epithelium of nasal polyps is thin, contains more mucocytes, and does not show the characteristic epithelial transmigration of neutrophils. In contrast to inverted papilloma, REAH occurs primarily on the posterior nasal septum rather than the lateral nasal wall and/or paranasal sinuses [2766]. REAH is also composed of numerous glands lined by respiratory epithelial cells, surrounded by thick hyalinized basement membranes, features not seen in inverted papilloma.
Invasive carcinoma can be distinguished from inverted papilloma by the presence of the following features: cellular pleomorphism, atypical mitoses, keratin pearls, loss of basement membranes, and stromal invasion associated with an inflammatory-desmoplastic response.

Prognosis and predictive factors
Though histologically benign, they have an unlimited growth potential and, if neglected, can cause considerable morbidity or even death by extending into contiguous structures. Attempts to remove these lesions intranasally by snare and avulsion have resulted in recurrence (or persistence) rates of 0-74% (average, 60%) (1442).

The preferred treatment for most lesions is a lateral rhinotomy and medial maxillectomy with meticulous removal of all mucosa in the ipsilateral paranasal sinuses. With this approach, the recurrence rate is usually <20% (1442). Selected small tumours can be effectively removed by a less aggressive approach using endoscopic sinonasal surgery. Recurrences typically appear within 2-3 years of therapy but, in some instances, are delayed for many years. Attempts to correlate histological features with risk of recurrence have resulted in conflicting data (1158,1442,2417). Even those with prominent mitotic activity and dysplasia do not invariably show an increased recurrence or malignancy. Nevertheless, dysplasia, especially if moderate to severe, demands thorough microscopic evaluation of all resected tissue to avoid overlooking small foci of carcinoma. The association between presence of HPV and the risk of recurrence is debatable (185,839).

Oncocytic papilloma (Schneiderian papilloma, oncocytic type)
Definition
A papilloma derived from the Schneiderian membrane composed of both exophytic fronds and endophytic invaginations lined by multiple layers of columnar cells with oncocytic features. Intraepithelial microcysts containing mucin and neutrophils are characteristic.

ICD-O code 8121/1

Synonyms
Oncocytic Schneiderian papilloma, cylindrical cell papilloma, columnar cell papilloma, papillomatosis.

Epidemiology
Oncocytic papilloma is equally distributed between the sexes, and the majority of the patients are aged over 50 years.

Etiology
In contrast to exophytic and inverted papillomas, HPV has not been identified in oncocytic papillomas (127).

Localization
Oncocytic papilloma almost always occurs unilaterally on the lateral nasal wall or in the paranasal sinuses, usually the maxillary or ethmoid. It may remain localized, involve both areas, or if neglected, extend into contiguous areas such as the orbit or cranial cavity.

Clinical features
Oncocytic papilloma presents as a fleshy, pink, tan, red-brown, or grey papillary or polypoid growth associated with nasal obstruction and intermittent epistaxis.

Histopathology
The oncocytic papilloma exhibits both exophytic and endophytic patterns of growth. The epithelium is multilayered, 2-8 cells thick, and is composed of tall columnar cells with swollen, finely granular cytoplasm reminiscent of oncocytes. The high content of cytochrome c oxidase and presence of numerous mitochondria ultrastructurally clearly establish their oncocytic character (129). The nuclei are either small dark and uniform or slightly vesicular with barely discernible nucleoli. Cilia in varying stages of regression may be observed in a few of the outermost cells.

The epithelium characteristically contains numerous small cysts filled with mucin or neutrophils (microabscesses). The stroma varies from edematous to fibrous, and may contain modest numbers of lymphocytes, plasma cells, and neutrophils, but few eosinophils. Seromucinous glands are sparse to absent.

Oncocytic papilloma and carcinoma
Four to 17% of all oncocytic papillomas harbour a carcinoma (1158,1266,1611,2723). Most of these are squamous, but mucoepidermoid, small cell and sinonasal undifferentiated carcinomas have also been described.

As in inverted papilloma, the carcinoma complicating oncocytic papilloma may arise within the papilloma, as evidenced by a gradation of histologic changes ranging from dysplasia to in situ to invasive carcinoma, or it may only be associated with the papilloma. Prognosis depends on the histologic type, degree of invasion, and the extent of tumour. In some instances, the carcinoma is in situ and of little consequence to the patient, whereas others are locally aggressive and may metastasize.

Differential diagnosis
The intraepithelial mucin-filled cysts of an oncocytic papilloma are often mistaken for rhinosporidiosis. In rhinosporidiosis, the organisms are not limited to the epithelium but also involve stroma, and do not induce a diffuse oncocytic change.

Oncocytic papilloma is also occasionally confused with a low-grade papillary ade-
The presence of intact basement membranes and absence of infiltrative growth are features that indicate a benign lesion. In addition, the presence of intraepithelial mucin-filled cysts and microabscesses and the stratified oncocytic epithelium of a papilloma are rarely seen in a low-grade adenocarcinoma.

**Prognosis and predictive factors**
The clinical behaviour parallels that of the inverted papilloma. If inadequately excised, at least 25-35% will recur, usually within 5 years. Smaller tumours can be resected endoscopically.

**Exophytic papilloma (Schneiderian papilloma, exophytic type)**

**Definition**
A papilloma derived from the Schneiderian membrane composed of papillary fronds with delicate fibrovascular cores covered by multiple layers of epithelial cells.

**ICD-O code** 8121/0

**Synonyms**
Fungiform papilloma, everted papilloma, transitional cell papilloma, septal papilloma, squamous papilloma, papillomatosis, Ringertz tumour

**Epidemiology**
Exophytic papillomas are 2-10 times more common in men, and occur in individuals between 20 and 50 years of age (2-87 years) [1158,1908].

**Etiology**
There is increasing evidence to suggest that exophytic papillomas may be etiologically related to HPV, especially types 6 and 11, rarely types 16 and 57b. In a collective review of exophytic papillomas evaluated for the presence of HPV by in situ hybridization and/or the polymerase chain reaction, about half of the cases were HPV positive [131].

**Localization**
Exophytic papillomas arise on the lower anterior nasal septum with no significant lateralization. As they enlarge, they may secondarily involve, but only infrequently originate from the lateral nasal wall.

**Clinical features**
Epistaxis, unilateral nasal obstruction, and the presence of asymptomatic mass are the typical presenting symptoms. On physical examination, they appear as papillary or warty, grey, pink or tan, non-translucent growths attached to the nasal septum by a relatively broad base.

**Histopathology**
Most exophytic papillomas range up to about 2 cm. Microscopically, they are composed of papillary fronds with fibrovascular cores covered by epithelium, 5-20 cells thick, that vary from squamous to transitional (intermediate) to ciliated pseudostratified columnar (respiratory). Scattered mucocytes are common. Surface keratinization is absent or scant, unless the lesion has been irritated or if the papilloma is unusually large and hangs into the nasal vestibule, where it is exposed to the drying effect of air. Mitoses are rare and never atypical. Unless infected or irritated, the stroma contains few inflammatory cells.

**Malignant change** in exophytic papilloma is exceptional [301,1908].

**Differential diagnosis**
Exophytic papillomas must be distinguished from the much more common, keratinizing cutaneous papillomas (e.g. verruca vulgaris) occurring in the nasal vestibule. The lack of extensive surface keratinization, presence of mucocytes, and presence of ciliated and/or ‘transitional’ epithelium help to confirm a diagnosis of exophytic papilloma. The presence of seromucinous glands and septal cartilage further indicate that the lesion is of mucosal rather than cutaneous origin.

**Prognosis and predictive factors**
Complete surgical excision is the treatment of choice. Inadequate excision rather than multiplicity of lesions probably accounts for the local recurrence of 22-50% [1158,1908].
**Respiratory epithelial adenomatoid hamartoma**

**Definition**
Benign nonneoplastic overgrowth of indigenous glands of the nasal cavity, paranasal sinuses and nasopharynx associated with the surface epithelium, and devoid of ectodermal neuroectodermal, and/or mesodermal elements.

**Synonyms**
Glandular hamartoma; seromucinous hamartoma.

**Epidemiology**
Hamartomas of the sinonasal tract and nasopharynx are uncommon. The majority of them are of pure epithelial type (respiratory epithelial adenomatoid hamartoma) [2766], although pure mesenchymal hamartomas or mixed epithelial-mesenchymal hamartomas may also rarely occur [14,106,933,2766]. Respiratory epithelial adenomatoid hamartomas predominantly occur in adult patients with a decided male predominance; patients range in age from the 3rd to 9th decades of life, with a median age in the 6th decade [2766].

**Etiology**
Respiratory epithelial adenomatoid hamartomas often arise in the setting of inflammatory polyps, raising a possible developmental induction secondary to the inflammatory process [2766].

**Localization**
The majority occur in the nasal cavity, in particular the posterior nasal septum; involvement of other intranasal sites occurs less often and may be identified along the lateral nasal wall, middle meatus and inferior turbinate [2766]. Other sites of involvement include the nasopharynx, ethmoid sinus, and frontal sinus. Most are unilateral, but some may be bilateral.

**Clinical features**
Patients present with nasal obstruction, nasal stuffiness, epistaxis and/or chronic (recurrent) rhinosinusitis. The symptoms may occur over months to years. Associated complaints include allergies.

**Macroscopy**
Lesions are typically polypoid or exophytic with a rubbery consistency, tan-white to red-brown appearance, measuring up to 6 cm in greatest dimension [933,2766].

**Histopathology**
The lesions are dominated by a glandular proliferation composed of widely-spaced, small to medium-sized glands separated by stromal tissue. In areas, the glands arise in direct continuity with the surface epithelium, which invaginate downward into the submucosa. The glands are round to oval, and composed of multilayered ciliated respiratory epithelium often with admixed mucocytes. Glands distended with mucus can be seen. A characteristic finding is stromal hyalinization with envelopment of glands by a thick, eosinophilic basement membrane. Atrophic glandular alterations may be present in which the glands are lined by a single layer of flattened to cuboidal epithelium. Small reactive seromucinous glands can be seen. The stroma is oedematous or fibrous, and contains a mixed chronic inflammatory cell infiltrate. Additional findings may include inflammatory sinonasal polyps, hyperplasia and/or squamous metaplasia of the surface epithelium unrelated to the adenomatoid proliferation, osseous metaplasia, rare association with inverted type Schneiderian papilloma, and rare association with a solitary fibrous tumour [2766].

**Prognosis and predictive factors**
Conservative but complete surgical excision is curative.

**Fig. 1.33** Respiratory epithelial adenomatoid hamartoma (REAH). A The glandular proliferation arises in direct continuity with the surface epithelium with invagination downward into the submucosa. Clusters of seromucinous glands are seen (arrow). B Pseudostratified epithelium with cilia within the adenomatoid collections of a REAH hamartoma. C Cilia along the luminal border of the cells (arrows). D Atrophic changes in which the glands are lined by a single layer of flattened to cuboidal-appearing epithelium. Note the prominent thickened stromal hyalinization enveloping the glands.
Among glandular tumours of the sinonasal tract, about one-quarter of cases are benign, and practically all of them are salivary gland-type neoplasms [1039]. For details see Chapter 5 on ‘Tumours of salivary glands’.

ICD-O codes
Pleomorphic adenoma 8940/0
Myoepithelioma 8982/0
Oncocytoma 8290/0

**Pleomorphic adenoma**
Most patients are between 20 and 60 years of age. Signs and symptoms are non-specific, and include unilateral nasal obstruction, epistaxis and a discernible mass. The tumour may resorb bone and extend into the maxillary sinuses. Most cases arise from the submucosa of the bony or cartilaginous nasal septum, but some arise in the lateral nasal wall [483,974,1210,1506]. The size varies from 0.5-5 cm [483] and tumours usually form polypoid, sessile swellings. Microscopically, they are unencapsulated, and tend to be cellular with predominance of modified myoepithelial cells often of plasmacytoid hyaline type; stromal elements are sparse. Exceptionally, focal skeletal muscle differentiation can occur [1419]. If treated by wide surgical excision, recurrence is uncommon [483].

**Myoepithelioma**
Myoepithelioma, including the spindle cell variant, of the sinonasal tract is very rare [188].

**Oncocytoma**
Oncocytomas of the sinonasal tract are rare, and most arise from the nasal septum [1039]. They are usually small, but some extend posteriorly and can cause bone resorption [470,480,998]. The nasolacrimal duct may be involved, causing unilateral epiphora and purulent rhinorrhoea [555]. Those examples that have behaved aggressively [470] are more appropriately considered low-grade oncocytic adenocarcinomas rather than adenomas [449,605,1044].
Fibrosarcoma

Definition
A malignant tumour of fibroblastic/myofibroblastic phenotype.

ICD-O code 8810/3

Synonyms
Fibromyxosarcoma; chondromyxofibrosarcoma.

Epidemiology
The incidence of sinonasal tract fibrosarcomas is difficult to determine because the diagnosis is often one of exclusion. These tumours are rare, accounting for <3% of all non-epithelial tumours. However, they are considered the second most common soft tissue sarcoma after rhabdomyosarcoma in the head and neck [168, 345, 826, 1041, 1317, 2438]. They occur in all ages, with a peak in the 5th decade. There is a 3:2 female:male gender predilection [168, 345, 826, 1041, 1317, 2438, 2511].

Etiology
A few patients have developed fibrosarcoma within the field of prior irradiation.

Localization
Most fibrosarcomas originate in one or more paranasal sinuses, while origination confined to the nasal cavity alone is less common [168, 345, 826, 1041, 1317, 2438]. The “infantile-type” fibrosarcoma in the sinonasal tract is exceedingly uncommon in the sinonasal tract and occurs near the choana [349, 1041, 1317].

Clinical features
Nearly all patients have nasal obstruction, often associated with epistaxis, while pain, sinusitis, nasal discharge, swelling, anosmia, and proptosis are less common. The median duration of symptoms is quite short.

Macroscopy
The tumours are smooth, nodular, pedunculated, fungating or ulcerating. The lesions range in size between 2 and 8 cm, with the cut surface revealing a circumscribed but not encapsulated, fleshy, homogeneous white-tan to yellow-pink mass, variably firm dependent upon the collagen content. Necrosis and haemorrhage may be present in higher-grade tumours.

Histopathology
The tumours are unencapsulated, sometimes sharply circumscribed, although often infiltrative and occasionally ulcerating. Bone invasion is common. Surface epithelial invagination into the tumour can be prominent, simulating an inverted papilloma. Spindle cells are arranged in compact fascicles, intersected by various amounts of delicate thin to dense keloid-like collagen. The cell bundles are arranged at acute angles to one another, occasionally giving rise to a “herringbone” or “chevron” pattern, while in most areas there is a more subtle fasciculation. A prominent storiform pattern is not seen. There is a marked variability in the cellularity within and between tumours. The cells are fusiform with a centrally placed hyperchromatic, needle-like nucleus surrounded by tapering cytoplasm which is often indistinct, creating a syncytial appearance to the fascicles. Most sinonasal tract fibrosarcomas are low grade. Nuclear pleomorphism is usually slight to moderate, but occasionally prominent. Mitotic figures are found in variable numbers. Haemorrhage and necrosis can be found in the poorly differentiated forms, with areas of myxoid degeneration. Focal osteo-cartilaginous differentiation has been described [168, 345, 826, 1041, 1317, 2438]. Fibrosarcomas are immunoreactive with vimentin, and sometimes focally with actin [1041].
Differential diagnosis
The differential diagnoses include malignant fibrous histiocytoma, spindle cell carcinoma, spindle malignant melanoma, malignant peripheral nerve sheath tumour, monophasic synovial sarcoma, rhabdomyosarcoma, glomangiopericytoma, desmoid fibromatosis, and nodular fasciitis [168,826,1041,2332].

Histogenesis
The (myo)fibroblast is considered the progenitor cell for these tumours.

Prognosis and predictive factors
Surgery is the treatment of choice, often followed by radiation therapy, yielding an overall long-term survival of 75% in low grade and localized tumours. The high incidence of recurrence (about 60%) is perhaps related to the complexity of the anatomy of the sinonasal tract and consequent difficulties of complete excision. Recurrence usually precedes metastasis, which occurs in about 15% of cases, most commonly to the lungs and bones and only rarely to lymph nodes. Poor prognostic factors include male gender, large tumour size, involvement of more than one contiguous site (nasal cavity and sinus, multiple sinuses), high histologic grade, and positive surgical margins [168,345,826,1041,1317,2438].

Malignant fibrous histiocytoma
Definition
Malignant fibrous histiocytoma (MFH) is currently used as a diagnosis of exclusion for sarcomas composed largely of myofibroblasts or undifferentiated mesenchymal cells.

ICD-O code 8830/3

Synonyms
Fibroxanthosarcoma, malignant fibrous xanthoma, myxofibrosarcoma, myxoid malignant fibrous histiocytoma

Epidemiology
Although once considered the most common sarcoma of adults, the frequency of its diagnosis has diminished since the introduction of immunohistochemistry has allowed assignment of some pleomorphic sarcomas to specific sarcoma entities. Only 3% of MFH occur in the head and neck, with 30% of these arising in the sinonasal area [2187]. MFH rarely occurs in the nasopharynx [1032,1923]. Sinonasal MFH most commonly occurs in adults with a male predominance [2433].

Etiology
Many sinonasal and nasopharyngeal MFH are a result of previous radiation, after a long latency period [1180,1345].

Localization
The maxillary sinus is most commonly affected, followed by the ethmoid sinuses and nasal cavity, whereas the frontal and sphenoid sinuses and nasopharynx are affected far less commonly [279,536,581,1032,1923,1936,2256,2426,2433].

Differential diagnosis
The differential diagnoses include fibrosarcoma, rhabdomyosarcoma, leiomyosarcoma, monophasic synovial sarcoma, malignant peripheral nerve sheath tumour, spindle cell carcinoma, spindle cell malignant melanoma and anaplastic large cell lymphoma.

Prognosis and predictive factors
Compared with other anatomical sites, MFHs of the head and neck generally have a slightly lower rate of recurrence and metastasis [133].

Macroscopy
Tumours are generally smooth, nodular or pedunculated (polypoid), with a number being fungating or ulcerating. The cut surface reveals a fleshy, homogeneous white-tan to yellow-pink mass with necrosis and haemorrhage, measuring up to 8 cm in maximum dimension [714].

Tumour spread and staging
Sinosal MFH can directly extend into nasopharynx, orbit, and pituitary fossa [1032,1707] and commonly metastasizes to lungs, bones, liver, [1707,2426] and only rarely to lymph nodes [279,1032,1707].

Histopathology
Sinosal MFH are generally infiltrative and ulcerative, but can occasionally be circumscribed. Pleomorphic MFH, the most frequent morphologic subtype of MFH in the sinonasal tract, is characterized by spindled to pleomorphic cells in a storiform growth pattern, with easily identified mitotic figures, including atypical forms, and necrosis. The cells are fusiform with tapering indistinct cytoplasm. Tumour giant cells with multiple nuclei may be found.

Immunohistochemistry
MFH are usually positive for vimentin and focally for actins. Importantly, MFH is a diagnosis of exclusion and is generally negative for desmin, skeletal muscle specific markers, S100 protein, HMB-45, epithelial markers and lymphoid markers.

Differential diagnosis
The differential diagnoses include fibrosarcoma, rhabdomyosarcoma, leiomyosarcoma, monophasic synovial sarcoma, malignant peripheral nerve sheath tumour, spindle cell carcinoma, spindle cell malignant melanoma and anaplastic large cell lymphoma.

Prognosis and predictive factors
Compared with other anatomical sites, MFHs of the head and neck generally have a slightly lower rate of recurrence and metastasis [133].

Fig. 1.38 Malignant fibrous histiocytoma. Remarkably pleomorphic cells with atypical mitotic figures.
Leiomyosarcoma

Definition
A malignant tumour of smooth muscle phenotype.

ICD-O code 8890/3

Epidemiology
Only a small number of sinonasal leiomyosarcomas have been reported {151, 824, 840, 1144, 1395, 1416, 1529, 2147, 2240, 2553}, accounting for <1% of all non-epithelial tumours. They occur in all ages, with a peak in the 6th decade (mean, 53 years) without a gender difference.

Etiology
There are a few reported cases with a prior history of radiation {824, 1416, 2147} or chemotherapy (cyclophosphamide specifically) {1416, 2147}.

Localization
Involvement of both the nasal cavity and paranasal sinuses is more common than involvement of the nasal cavity alone {824, 840, 1144, 1395, 1416, 1529, 1745}.

Clinical features
Nearly all patients have nasal obstruction, frequently associated with epistaxis and pain, while nasal discharge, swelling, and blurred vision are less common. The duration of symptoms is usually long {824, 840, 1144, 1395, 1416, 1529, 2147, 2240, 2553}. There is usually no lymphadenopathy. Plain radiographs show opacification of the nasal cavity or sinus(es), often suggesting sinusitis {1144, 1395, 1529, 2553}.

Macroscopy
These tumours range in size up to 7 cm, with an average of about 4 cm. They are more likely infiltrative than circumscribed, and occasionally polypoid. The surface is typically ulcerated and crusted. These bulky tumours have a cut surface which reveals a soft to firm, grey-white and fleshy appearance. Haemorrhage, necrosis and cystic change are common.

Histopathology
Leiomyosarcomas are infiltrative neoplasms accompanied by surface ulceration. Bone or cartilage invasion is more frequent than surface or seromucinous gland invasion. Leiomyosarcomas are composed of right-angle intersecting bundles of spindle cells. Palisading, storiform and “haemangiopericytoma-like” patterns can occur. The tumours are hypercellular, but coagulative tumour necrosis and haemorrhage can create a hypocellular appearance. The tumour cells have elongated, vesicular to hyperchromatic, lobulated or indented nuclei with blunt ends (“cigar-shaped”). The cytoplasm is fibrillary and eosinophilic, with frequent perinuclear vacuolation. Mitoses, both typical and atypical, are present to a variable degree {824, 840, 1144, 1395, 1416, 1529, 2147, 2240, 2553}.

Electron microscopy
Electron microscopy reveals variable features of smooth muscle cells, including myofilaments arranged in parallel arrays, dense bodies within the filaments, cell junctions, pinocytotic vesicles and basal lamina {1395, 1529, 1933}.

Differential diagnosis
The differential diagnoses include sinonasal glomangiopericytoma, peripheral...
eral nerve sheath tumour, fibrosarcoma, spindle cell carcinoma and melanoma (824,840,1144,1395,1416,1529,2147,2240,2553,2603).

**Genetic susceptibility**
There are isolated cases of children with leiomyosarcomas who have preexisting hereditary retinoblastomas (627).

**Prognosis and predictive factors**
About half of the reported cases develop local recurrence, often within one year, and nearly 1/3 of these patients will subsequently develop metastasis (mostly to the lungs and liver). Complete surgical excision is difficult to achieve, and radiation and chemotherapy are used with variable results (824,1416,2501). Poor prognostic factors include involvement of more than one contiguous site, large tumour size (>5 cm), high mitotic count (>20/10 high power field), tumour necrosis, and tumour stage (824,840,1144,1395,1416,1529,2147,2240,2553).

**Rhabdomyosarcoma**

**Definition**
A malignant tumour of skeletal muscle phenotype.

**ICD-O code** 8900/3
(Also see subtypes: 8910/3, 8912/3, 8920/3, 8901/3 in WHO Tumours of Soft Tissue)

**Synonyms**
Myosarcoma, malignant rhabdomyoma, rhabdosarcoma, embryonal sarcoma, rhabdomyoblastoma

**Epidemiology**
Approximately 40% of rhabdomyosarcomas occur in the head and neck (1978), with about 20% in the nasal cavity, nasopharynx, and nasal sinuses (2745). Rhabdomyosarcoma is the most common sarcoma in childhood. The embryonal subtype predominates in children, while the alveolar subtype predominates in adults (825,1273). The pleomorphic subtype is rare (836,837). There is an overall slight male predominance (825).

**Localization**
The nasopharynx is more commonly involved than the sinonasal tract (326,724). In adults, rhabdomyosarcoma is more common in the ethmoid sinuses, followed by the maxillary sinuses and nasopharynx (1856).

**Clinical features**
Signs and symptoms include difficulty in breathing, epistaxis, facial swelling, visual disturbances, and sinusitis often of short duration. Tumours may appear as a large, polypoid sinonasal mass or may occasionally protrude as a gelatinous mass from the nares (825). CT and MRI imaging delineate the size and extent of the tumour (1453,2846). The botryoid type shows grape-like rings and heterogeneous enhancement (980).

**Macroscopy**
The embryonal subtype is generally poorly circumscribed, fleshy, pale and tan; the spindle cell variant is firm, fibrous, and tan-yellow with a whorled cut surface. The botryoid variant has a grape-like or polypoid appearance (825). The alveolar subtype is fleshy to firm tan-grey.

**Tumour spread and staging**
These tumours often spread to contiguous sites including base of the skull, temporal bones, and orbit (724,825). About 40% metastasize to lymph nodes, bones, and lungs, and less commonly bone marrow, soft tissue, liver and brain (1441,1856). The tumours are staged according to the Intergroup Rhabdomyosarcoma Study. Group I includes local disease, Group II residual disease or local spread, Group III incomplete resection or biopsy with gross residual disease, and Group IV metastatic disease at onset (613). Most adult sinonasal and nasopharyngeal rhabdomyosarcomas are staged as Group III or IV at presentation (1856).

**Histopathology**
Embryonal rhabdomyosarcoma has round to spindled cells with hyperchromatic nuclei. Larger rhabdomyoblasts with eosinophilic cytoplasm are usually identified, but cross striations are difficult to recognize. Myxoid stroma is common. The spindle cell variant, characterized by spindled cells in fascicular to storiform growth patterns, can be deceptively bland. The botryoid variant is polypoid with a submucosal hypercellular cambium layer, a myxoid hypocellular zone, and a deep cellular component. Alveolar rhabdomyosarcoma typically has fibrous septa separating clusters of loosely cohesive groups of small to medium round tumour cells with hyperchromatic nuclei and scant eosinophilic cyto-

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Fig. 1.41 Rhabdomyosarcoma  A Nasal alveolar rhabdomyosarcoma, with typical alveolar pattern.  B The tumour cells are ovoid and have hyperchromatic nuclei and scant eosinophilic cytoplasm.
Malignant soft tissue tumours

plasm. Multinucleated giant cells with overlapping peripheral nuclei are often present. The solid variant grows in sheets and lacks septa. Rarely, the tumour can be composed exclusively or predominantly of clear cells. A mixed alveolar and embryonal pattern may occur. Mitotic figures are usually easy to identify. Pleomorphism is occasionally observed focally. After treatment, there is often increased cytodifferentiation, with the cells exhibiting abundant eosinophilic fibrillary cytoplasm (2644). Pleomorphic rhabdomyosarcoma is rare and uncommon in this location.

**Immunohistochemistry**

There is immunoreactivity for desmin, muscle specific actin, myoglobin, fast myosin, nuclear MyoD1 and nuclear myogenin (skeletal muscle myogenin, myf4) (2619). CD99 can be positive in 16% of cases (1084).

**Electron microscopy**

Electron microscopy shows some degree of skeletal muscle differentiation ranging from well-formed Z-bands to incomplete sarcomeres with thick and thin filaments and ribosome-myosin complexes (724,837).

**Differential diagnosis**

The differential diagnoses of embryonal rhabdomyosarcoma include sinonasal polyp with stromal atypia (1840) and various sarcomas. The differential diagnoses of alveolar rhabdomyosarcoma include various round blue cell tumours, including lymphoma, sinonasal undifferentiated carcinoma, small cell carcinoma of neuroendocrine type, mesenchymal chondrosarcoma, PNET/Ewing sarcoma, olfactory neuroblastoma, and mucosal malignant melanoma.

**Somatic genetics**

Embryonal rhabdomyosarcoma shows allelic loss at 11p15 (271,925). Alveolar rhabdomyosarcoma has a consistent translocation, usually t(2;13) (PAX3-FKHR), or less commonly t(1;13) (PAX7-FKHR) which can be performed on paraffin-embedded sections (141).

**Genetic susceptibility**

Germline mutations of TP53 in Li-Fraumeni syndrome are found in some children with rhabdomyosarcoma.

**Prognosis and predictive factors**

Prognosis is determined by patient age, histologic subtype, and tumour clinical group (2123). Younger patients have a more favourable prognosis than older

![Fig. 1.42 Sinonasal rhabdomyosarcoma. A This macroscopic image demonstrates the similarity between a sinonasal polyp and a rhabdomyosarcoma. B Irregular islands of tumour in stroma may mimic carcinoma. Desmin and skeletal muscle markers can aid in the diagnosis of rhabdomyosarcoma.](image)

![Fig. 1.43 A Alveolar rhabdomyosarcoma of nasal cavity. After chemotherapy, there is increased cytodifferentiation to rhabdomyoblasts. B An embryonal rhabdomyosarcoma demonstrates remarkably atypical spindle cells with small amounts of eosinophilic cytoplasm.](image)
patients in all rhabdomyosarcoma subtypes. Currently, the 5-year survival is 44-69%, and 90% for clinical Group I disease [322,1084]. Adults have a poor prognosis, with 5-year survival of <10% [1841,1856]. Embryonal rhabdomyosarcoma has a better prognosis than alveolar rhabdomyosarcoma [2123]. Botryoid and spindle cell variants [346] have a better prognosis than embryonal rhabdomyosarcoma. Furthermore, alveolar rhabdomyosarcomas with PAX7/FKHR are thought to have better prognosis than PAX3/FKHR tumours [1298].

Angiosarcoma

Definition
A malignant neoplasm of vascular phenotype whose constituent tumour cells have endothelial features.

ICD-O code 9120/3

Synonyms
Malignant haemangioendothelioma; malignant angioendothelioma; lymphangiosarcoma, haemangiosarcoma.

Epidemiology
Angiosarcoma is uncommon, accounting for less than 1% of all sinonasal tract malignancies [89,1603,1640,1848]. They occur in all ages, with a peak in the 5th decade, and a male predilection (male:female = 2:1). Females tend to be younger at presentation by up to a decade [823,1848,2633,2795,2812].

Etiology
Radiation exposure [1556,1603,1848], Thorotrast, arsenic and vinyl chloride are reported risk factors [2795].

Localization
The maxillary sinus is most frequently affected. Other sites that may be involved primarily or secondarily include the nasal cavity and other paranasal sinuses [823,1848,2633,2795,2812].

Clinical features
Patients present with recurrent epistaxis, profound pallor, a mass lesion, pain (including headache, otalgia, toothache), nasal obstruction, sinusitis, nasal discharge (often described as foul-smelling and blood tinged), paraesthesia and/or loose teeth. The duration of symptoms ranges from weeks to months, but is generally short (median, 4 months).

Macroscopy
The tumours range up to 8 cm, with a mean of about 4 cm. They are nodular, polypoid and morulated, soft and friable, purple to red, often ulcerated with associated haemorrhage or clot and necrosis [823,1848,2633,2795,2812].

Histopathology
Most sinonasal angiosarcomas are histologically low-grade. They infiltrate the adjacent tissues and bone, accompanied by necrosis and haemorrhage. They comprise tortuous anastomosing vascular channels that dissect the stroma, capillary-sized vessels and cavernous vascular spaces. The lining endothelial cells range from flat to plump spindly to epithelioid, and often form papillary tufts. Intracytoplasmic vacuoles (neolumen), often containing erythrocytes, are characteristic of the epithelioid variant. The degree of nuclear pleomorphism is variable. Mitotic figures, including atypical forms, are variably present [823,1848,2633,2795,2812].

Immunohistochemistry
Angiosarcomas are immunoreactive for CD34, CD31, Factor VIII R-Ag and vimentin, and focally keratin (especially the epithelioid variant) and actin [2812].

Differential diagnosis
The differential diagnoses include granulation tissue, intravascular papillary
endothelial hyperplasia, haemangioma, nasopharyngeal angiofibroma, angiolymphoid hyperplasia with eosinophilia, glomangiopericytoma, Kaposi sarcoma, malignant melanoma, carcinoma and large cell lymphoma [30,1388,1976, 2469,2764].

**Malignant soft tissue tumours**

**Prognosis and predictive factors**

Patients are usually treated by surgical resection with radiation and/or chemotherapy. Recurrences are common (50%), likely due to incomplete excision or possible multifocality. Metastasis is uncommon, and the predilection sites are the lung, liver, spleen, and bone marrow [1976]. The outcome is more favourable compared with the almost uniformly fatal outcome for cutaneous and soft tissue angiosarcomas [823, 1848,2633,2795,2812]

**Malignant peripheral nerve sheath tumour**

**Definition**

A malignant tumour of nerve sheath phenotype.

**ICD-O code**

9540/3

**Synonyms**

Neurogenic sarcoma, malignant schwannoma, neurofibrosarcoma.

**Epidemiology**

Malignant peripheral nerve sheath tumours (MPNSTs) comprise 2-14% of all head and neck sarcomas [1231,1562], arising de novo or less commonly in the setting of neurofibromatosis type 1 (NF1) [1059,1231,1795]. De novo MPNST peaks in the fourth decade, while those in the setting of NF1 occur at an earlier age. There is a female predominance for de novo sinonasal MPNST [1041], and a male predominance in NF1-associated MPNST [774].

**Etiology**

Radiation and possibly immunosuppression may be etiologic factors [1562].

**Localization**

They commonly arise from the ophthalmic and maxillary branches of the trigeminal (5th) cranial nerve, but can involve all of the sinonasal tract and nasopharynx [756,1153,1795,2018].

**Clinical features**

Presenting symptoms include mass, pain, epistaxis, deviation or swelling of tonsils, nasal obstruction, and sinusitis [27,1319,1891,1964,2119].

**Macroscopy**

MPNST is generally globoid to fusiform, pseudoencapsulated, cream-grey and firm, occasionally associated with surface ulceration. Infiltration into the surrounding soft tissues and bone is common. The tumours are often large (>5 cm) and may be attached to a nerve. Foci of cyst formation, necrosis and/or haemorrhage are frequent.

**Tumour spread and staging**

Local extension into contiguous structures along the path of the trigeminal nerve or through the foramen ovale are characteristic [1452]. MPNSTs metastasize to the lungs, bones, and/or liver, [1041] while the epithelioid variant tends to involve regional lymph nodes [1437].

**Histopathology**

MPNSTs can either be spindled (95%) or epithelioid (5%) [1437]. At low magnification, both types show alternating areas of dense cellularity with less cellular myxoid areas. Geographic necrosis and perivascular accentuation of tumour cells are common. The tumour cells are fusiform and plump, arranged in tightly packed fascicules woven into a vague “herringbone” pattern, while in other areas the cells are wavy with fibrillar cytoplasmic extensions, arranged in a loose myxoid.
Tumours of the nasal cavity and paranasal sinuses

background matrix. Focal palisading of nuclei may be present. The tumour cells are variably pleomorphic, with a high nuclear to cytoplasmic ratio and mitotic activity. Many sinonasal tract MPNSTs, in contrast to those occurring in other anatomic sites, are histologically and biologically low-grade [1041]. An origin from a nerve may or may not be apparent. MPNST with rhabdomyoblasts are known as malignant Triton tumours.

Immunohistochemistry

The spindle cell variant is usually focally positive for S100 protein and occasionally positive for glial fibrillary acidic protein (GFAP). However, up to 30% of MPNST may be negative for S100 protein [2784A]. The epithelioid variant is diffusely immunoreactive for S100 protein and may mimic malignant melanoma, [583,756,2603] but other melanoma markers are negative. In malignant Triton tumour, the rhabdomyoblasts are positive for desmin and other skeletal muscle markers.

Differential diagnosis

The differential diagnoses include synovial sarcoma, fibrosarcoma, spindle cell carcinoma, leiomyosarcoma and mucosal malignant melanoma [1041,2550,2603].

Precursor lesions

MPNST may arise from neurofibroma (especially in the setting of NF1) and only exceptionally from classic schwannoma.

Somatic genetics

Both NF1 alleles must be inactivated for MPNST to occur in NF1. Malignant progression from neurofibroma in NF1 is related to alterations of genes controlling cell cycle regulation, including TP53 [1459] and CDKN2A (which encodes p16) [1361,1895].

Genetic susceptibility

MPNST of the sinonasal tract may be associated with NF1, typified by germline mutation of the NF1 tumour suppressor gene located on chromosome 17 [454].

Prognosis and predictive factors

Surgery is the treatment of choice, although radiation and chemotherapy may have a palliative role. De novo sinonasal MPNSTs have a 5-year survival rate of about 90%, which is superior to that of 50-65% for MPNSTs arising in other anatomic locations [1041,1562,2715]. However, NF1-associated sinonasal MPNSTs have a 5-year survival rate of only about 15% [2119]. Poor prognostic factors include male gender, high tumour cellularity and high mitotic activity [1041].
Desmoid-type fibromatosis

Definition
A locally aggressive, cytologically bland tumour of (myo)fibroblastic phenotype.

ICD-O code 8821/1

Synonyms
Extra-abdominal desmoid, extra-abdominal fibromatosis, desmoid tumour, aggressive fibromatosis, juvenile desmoid-type fibromatosis, infantile fibromatosis.

Epidemiology
Although 15% of cases of desmoid-type fibromatosis occur in the head and neck, the sinonasal tract is uncommonly involved [6,903,2643]. All ages can be affected, especially children [903]. There is a male predilection [903].

Localization
The maxillary sinus and turbinates are usually affected, and the involvement can occasionally be bilateral [514,826,903].

Clinical features
Symptoms include nasal obstruction, epistaxis, mass, facial pain, tooth displacement, and a non-healing tooth extraction site [514,826,903].

Macroscopy
The lesion is tan-white, glistening, and rubbery to firm, and is often infiltrative. It measures up to 7 cm. Fibromatosis may be multicentric, especially in the setting of Gardner syndrome [562].

Histopathology
This is an infiltrative growth with low to moderate cellularity, comprising broad fascicles of bland-looking spindle cells and collagen fibers often arranged in a uniform direction. Elongated blood vessels are frequently observed running parallel to each other. The spindle cells have a myofibroblastic appearance, with low nuclear to cytoplasmic ratio and uniformly bland ovoid nuclei with indistinct nucleoli. Mitotic figures are infrequent and never atypical. The matrix is collagenized to focally myxoid, and keloid-like collagen may be present. The main differential diagnoses include hypertrophic scar and fibrosarcoma.

Genetic susceptibility
Fibromatosis can be part of Gardner syndrome [562].

Prognosis and predictive factors
Fibromatosis can be locally aggressive and involve contiguous structures, with approximately 20% recurrence rate, but it does not metastasize [514,903,2643]. Recurrence generally occurs within the first few years and is related to inadequacy of surgical margins [514,903,2643].

Inflammatory myofibroblastic tumour

ICD-O code 8825/1

Inflammatory myofibroblastic tumour uncommonly occurs in the sinonasal tract [2429]. Please see corresponding section in ‘Tumours of the hypopharynx, larynx and trachea’.

Glomangiopericytoma (Sinonasal-type haemangiopericytoma)

Definition
A sinonasal tumour demonstrating perivascular myoid phenotype.

ICD-O code 9150/1

Fig. 1.50 Desmoid-type fibromatosis. Heavily collagenized stroma with spindle cells with bland nuclei and elongated vessels.

Fig. 1.51 Glomangiopericytoma. Characteristic diffuse growth within the submucosa, with effacement of the normal components of the submucosa and preservation of mucoserous glands. The overlying respiratory epithelium remains intact.
Tumours of the nasal cavity and paranasal sinuses

Synonyms
Sinonasal haemangiopericytoma; haemangiopericytoma-like tumour, sinonasal glomus tumour; haemangiopericytoma.

Epidemiology
Sinonasal glomangiopericytomas predilect to the nasal cavity and paranasal sinuses, where they comprise <0.5% of all neoplasms [343,482,2600]. There is a very slight female predominance. All ages can be affected (in-utero to 86 years), but the peak is in the 7th decade.

Localization
Tumours most frequently arise unilaterally in the nasal cavity alone, although extension into paranasal sinuses can occur. Isolated paranasal sinus involvement is uncommon. Rarely, large tumours may appear to arise bilaterally [216,343,482,638,649,2600].

Clinical features
The majority of patients present with nasal obstruction, epistaxis, or non-specific findings, such as a mass, polyp, difficulty breathing, sinusitis, headache and nasal congestion, present for an average duration of <1 year. Imaging studies show nasal cavity or paranasal sinus opacification by a polypoid mass lesion, frequently accompanied by sinusitis, bone erosion and sclerosis [482,2600,2729].

Macroscopy
The generally polypoid tumours range up to 8 cm, with a mean size of about 3 cm. The tumours are beefy red to greyish pink, soft, edematous, fleshy to friable masses, often demonstrating haemorrhage.

Histopathology
This is a subepithelial well-delineated but unencapsulated cellular tumour, effacing or surrounding the normal structures. It is comprised of closely packed cells, forming short fascicles and sometimes exhibiting a storiform, whorled or palisaded pattern, interspersed with many vascular channels. The latter are in the form of capillary-sized to large patulous spaces that may have a “staghorn” or “antler-like” configuration. A prominent peritheliomatous hyalinization is characteristic. The neoplastic cells are uniform, elongated to oval, and possess vesicular to hyperchromatic, round to oval to spindle-shaped nuclei, and lightly eosinophilic cytoplasm. Mild nuclear pleomorphism and occasional mitotic figures may be present, but necrosis is not found. Extravasated erythrocytes, mast cells, and eosinophils are nearly ubiquitously present. Occasionally, tumour giant cells, fibrosis or myxoid degeneration may be seen.

Immunohistochemistry
Immunohistochemically, glomangiopericytoma is distinctly different from soft tissue haemangiopericytoma by yielding diffuse reactivity for actins, factor XIIIa and vimentin, and lacking strong diffuse staining for CD34. Bcl-2, FVIII-R Ag, CD99 and CD117 are negative [343,638,1364,2070,2600].

Differential diagnosis
The differential diagnoses include haemangioma, solitary fibrous tumour, glomus tumour, leiomyoma, synovial sarcoma and leiomyosarcoma.

Histogenesis
This tumour has been known as haemangiopericytoma-like tumour or sinonasal haemangiopericytoma, but it is clinically, morphologically and biologically distinct from soft tissue-type or dura-based haemangiopericytoma [446,544,773,936,1723,1724,2276,2386,2689]. The proposed cell of origin is a modified perivascular glomus-like myoid cell.

Prognosis and predictive factors
Sinonasal glomangiopericytoma is indolent, with an overall excellent survival (>90% 5-year survival) achieved with complete surgical excision. Recurrence, which develops in up to 30% of cases, may occur many years after the initial surgery [216,343,638,649,2600]. Aggressive-behaving glomangiopericytomas (malignant glomangiopericytomas) are uncommon [216,343,482,556,1779,2600], and usually exhibit the following features: large size (>5 cm), bone invasion, profound nuclear pleomorphism, increased mitotic activity (>4/10 high power fields), necrosis, and proliferation index >10% [216,343,1364,2600].
Extrapleural solitary fibrous tumour

ICD-O code 8815/1
Solitary fibrous tumours are tumours of CD34-positive fibroblasts which often show a prominent haemangiopericytoma-like vascular pattern. They are exceedingly uncommon in the upper respiratory tract, where they comprise <0.1% of all neoplasms. All ages can be affected without a gender predilection. Tumours can affect the nasal cavity, nasopharynx or paranasal sinuses. Patients present with nasal obstruction, epistaxis or other non-specific symptoms. The tumour is usually polypoid and firm. It is composed of a variably cellular proliferation of bland spindle-shaped cells with nondescript growth pattern associated with “ropy” keloidal collagen bundles and interlaced thin-walled vascular spaces. The latter may be prominent and exhibit a haemangiopericytoma-like pattern. Solitary fibrous tumours are immunoreactive for CD34 and bcl-2, and generally lack actin immunoreactivity. The diagnosis rests on a combination of architectural, cytomorphic, and immunophenotypic features. The differential diagnoses include sinonasal glomangiopericytoma, fibrous histiocytoma, leiomyoma, schwannoma, synovial sarcoma, and fibrosarcoma. Complete surgical removal yields the best patient outcome. Occasional cases may potentially show a malignant behaviour (158,834,997,1706,2600,2800, 2914). See WHO Classification of Tumours of Soft Tissue and Bone [775].

Fig. 1.53 Solitary fibrous tumour of nasal cavity. A Circumscribed tumour beneath epithelium. B Bland-looking spindly cells are tightly intertwined with collagen fibers.

Fig. 1.54 Solitary fibrous tumour A Cytologically bland cells are arranged in streaming fashion, with collagen deposited between the cells. Haemorrhage is seen. B A solitary fibrous tumour strongly and diffusely immunoreactive for CD34.
Benign soft tissue tumours

Myxoma

Myxoma is a benign soft tissue tumour characterized by bland spindle shaped cells embedded in hypovascular, myxoid stroma. For details see Chapter 6 (Odontogenic tumours).

Leiomyoma

Definition
A benign tumour of smooth muscle phenotype.

ICD-O code 8890/0

Synonyms
Angioleiomyoma; vascular leiomyoma; leiomyoblastoma

Epidemiology
Primary leiomyomas of the sinonasal tract are very rare (824,1144,1307, 1535,1796,2114,2635,2695). There is a peak in the 6th decade, although men are younger than women by a decade at initial diagnosis. There is a female predilection, with a ratio of 3.5:1.

Etiology
Other than prior radiation, there are no known risk factors.

Localization
The turbinates are affected most frequently, (824,1144,1307,1535,1796, 2114,2635,2695) with isolated cases reported in the paranasal sinuses alone or in combination with the nasal cavity (1842).

Clinical features
Nearly all patients have nasal obstruction, although nasal discharge, epistaxis, headaches and pain are also common (824,2114,2635,2695).

Macroscopy
These tumours have an average size of 2 cm, but rare ones may be as large as 10 cm. They are sessile or polypoid, with a smooth, well circumscribed border.

Histopathology
Leiomyomas are located in the submucosa, separated from a typically intact mucosa. They are composed of spindled cells arranged in orderly fascicles, whorls and intersecting bundles. The cells have elongated, vesicular to stippled nuclei with blunt ends (“cigar-shaped”), surrounded by spindled, bipolar, fibrillar eosinophilic cytoplasm. They are highly differentiated, with little or no atypia, although rare cells may exhibit nuclear pleomorphism (2695). Necrosis and invasion are absent, and mitotic activity is scarce. Mucinous degeneration, hyalinization or fibrosis, and adipocytes can be seen, but these features are usually focal and more likely seen in larger lesions (1144,1535,1796, 2695). Vascular leiomyoma (angiomyoma) contains capillary, cavernous or venous vascular spaces, with the smooth muscle cells being associated with the vessel walls and represents the most common type of benign smooth muscle tumour in this region.

Immunoprofile
The tumour cells are diffusely and strongly immunoreactive for actins, desmin, h-caldesmon and vimentin. The Ki-67 index is usually <5% (1144).

Differential diagnosis
The differential diagnoses include sinonasal glomangiopericytoma, haemangioma, peripheral nerve sheath tumour and leiomyosarcoma.

Prognosis and predictive factors
Complete excision is curative.

Haemangioma

Definition
A benign neoplasm of vascular phenotype.

Fig. 1.55 A Myxoma. The stellate cells have thin processes which extend out into the background mucinous matrix. B Vascular leiomyoma. Spindle tumour cells are identified scrolling off thick muscle-walled vessels.
ICD-O code 9120/0

Synonyms
Lobular capillary haemangioma; pyogenic granuloma; capillary haemangioma; cavernous haemangioma; epulis gravidarum.

Epidemiology
Mucosal haemangiomas of the nasal cavity, paranasal sinuses and nasopharynx account for 10% of all head and neck haemangiomas and approximately 25% of all non-epithelial neoplasms of this anatomical region. The haemangiomas occur in all ages, although there is a peak in children and adolescent males, females in the reproductive years, and then an equal distribution beyond 40 years of age. Patients with cavernous haemangiomas tend to be men in the 5th decade {166,167,658,823,1037,1189,1270,1738,2056,2333}.

Etiology
Lobular capillary haemangioma (pyogenic granuloma) has an association with injury and hormonal factors (pregnancy or oral contraceptive use) {2158}.

Localization
The septum is most frequently affected (specifically, the anterior septum in Little’s area), followed by the turbinates (usually the tip) and the sinuses {658,823,1037,1189,1270,1738,2056,2333}.

Clinical features
Patients present with unilateral epistaxis and/or an obstructive painless mass. Sinus lesions present as sinusitis, proptosis, mass, anaesthesia or pain. Symptoms are usually present for a short duration {658,823,1037,1189,1270,1738,2056,2333,2753}.

Macroscopy
The tumours range up to 5 cm, with a mean size of <1.5 cm. Grossly, they appear as a red to blue submucosal soft, compressible, flat or polypoid lesion, often with an ulcerated surface. Cavernous haemangiomas are spongy on sectioning {658,823,1037,1189,1270,1738,2056,2333,2753}.

Histopathology
Haemangiomas are usually localized and can be divided into capillary and cavernous types based on the size of the blood vessels. Haemangiomatosis is a more diffuse lesion often involving contiguous structures {823,1270,2126}. Lobular capillary haemangioma is a circumscribed lesion comprising lobules of capillaries lined by plump endothelial cells and supported by prominent pericytes. The lobules are separated by a fibromyxoid stroma. The cellularity of the lobules may be quite high. Mitotic figures are often observed, but are never atypical. The surface epithelium often forms collarettes around the lesion {658,823,1037,1189,1270,1738,2056,2333}. If the lesion is ulcerated and inflamed, the term ‘pyogenic granuloma’ has been applied. Cavernous haemangiomas are frequently intraosseous or involve the turbinates or lateral nasal wall. They are composed of multiple, large thin-walled, dilated blood vessels separated by scant fibrous stroma {658,823,1037,1189,1270,1738,2056,2333}.

Immunoprofile
The tumour cells are immunoreactive for Factor VIII related antigen, CD34, CD31 and Ulex europaeus I lectin. The proliferated blood vessels are enwrapped by actin-positive pericytes.

Differential diagnosis
Haemangiomas should be distinguished from granulation tissue, telangiectasia, vascular malformations, vascular polyps (haemorrhagic type), papillary endothelial hyperplasia, angiofibroma, bacillary angiomatosis, angiolymphoid hyperplasia with eosinophilia, glomus tumour, sinonasal glomangiopericytoma, lymphangioma, Kaposi sarcoma, and angiosarcoma. Haemangioma can be distinguished from granulation tissue by the lobular arrangement of the capillaries in the former and the more parallel arrangement of vessels in the latter. The distinction between a haemangioma and telangiectasia may be difficult but is facilitated in a patient with a known family history of hereditary haemorrhagic telangiectasia (Osler-Weber-Rendu syndrome) {658,823,1037,1189,1270,1738,2056,2073,2333,2600}.

Prognosis and predictive factors
Haemangiomas are generally easy to remove, although larger tumours may be complicated by excessive bleeding. They should be removed in all ages, especially in children since aplasia of the nasal cartilages may cause eventual disfigurement. If the tumour is pregnancy-
related, regression will often occur after parturition. Multiple recurrences are more common in children if the lesional bed is not completely eradicated.

**Schwannoma**

**Definition**
A usually encapsulated, benign tumour composed of differentiated, neoplastic Schwann cells.

**ICD-O code** 9560/0

**Synonyms**
Neurilemoma, neurilemmoma, benign peripheral nerve sheath tumour.

**Epidemiology**
Less than 4% of schwannomas involve the nasal cavity and paranasal sinuses [1091,2351], and they occur in middle-aged adults with an equal gender distribution [2351].

**Localization**
Sinonasal schwannomas arise from the branches of the trigeminal (5th) nerve and autonomic nervous system, and most commonly involve the ethmoid and maxillary sinuses, followed by the nasal cavity, sphenoid and frontal sinuses [1023,1091,2018,2351]. Cellular schwannoma tends to be located in the midline.

**Clinical features**
The presenting symptoms include obstruction, rhinorrhea, epistaxis, anosmia, headache, dysphagia, hearing loss, facial or orbital swelling, and pain [2018,2351].

**Macroscopy**
Sinonasal schwannoma ranges in size up to 7 cm. It is a well-delineated but non-encapsulated globular, firm to rubbery yellow-tan mass. The cut surfaces show tan-grey, yellowish, solid to myxoid and cystic tissue, commonly with haemorrhage.

**Tumour spread and staging**
The tumour can expand into the orbit, nasopharynx, pterygomaxillary fossa and cranial cavity [2351].

**Histopathology**
Schwannoma is composed of cellular Antoni A areas with Verocay bodies and hypocellular myxoid Antoni B areas. The cells are fusiform with elongated fibrillary cytoplasm, and buckled to spindled nuclei which show little pleomorphism, although scattered large pleomorphic or bizarre cells can be present in some cases. Nuclear palisading is often evident in some foci. There are frequently small to medium-sized vessels with ectasia, thrombosis and perivascular hyalinization in the Antoni B areas. Extensive degenerative changes can occur, and may result in only a thin rim of recognizable tumour. Cellular variants exhibit only the Antoni A pattern, but no fascicular growth or Verocay bodies.

**Immunoprofile**
The tumour cells are strongly and diffusely immunoreactive for S100 protein. CD34 only stains some more slender cells in the Antoni B areas. Neurofilament is absent. GFAP and keratins may be positive.

**Prognosis and predictive factors**
Schwannoma is a benign tumour with a very low recurrence potential. Malignant transformation is exceptional [1690].

**Neurofibroma**

**Definition**
A benign tumour of peripheral nerve sheath phenotype with mixed cellular components, including Schwann cells, perineurial hybrid cells and intraneural fibroblasts.

**ICD-O code** 9540/0

**Epidemiology**
Neurofibromas are extremely rare in the sinonasal tract. In NF1-related neurofibromas, patients tend to be younger, with a male predominance [2745]. For the more common sporadic neurofibromas,
all ages may be affected, although patients tend to be older with an equal gender distribution.

**Localization**
The tumour arises from the ophthalmic or maxillary branches of the trigeminal (5th) nerve and is most commonly located in the maxillary and ethmoid sinuses and/or nasal cavity [288]. Plexiform neurofibroma may occur in the sinonasal area where it is found in the maxillary sinus [817], usually associated with NF1.

**Clinical features**
Symptoms include epistaxis, rhinorrhoea, swelling, mass, obstruction, and pain [61,2018].

**Macroscopy**
The tumour is firm, glistening, grey-tan, fusiform, and sometimes polypoid, in a submucosal location with an intact surface epithelium [61,1095].

**Histopathology**
Neurofibromas are generally submucosal paucicellular lesions. They are composed of spindled cells with wavy, dark-staining nuclei and scanty cytoplasm, in a background of wavy collagen fibres, myxoid stroma and mast cells. The centre of the lesion usually shows residual neurites.

**Immunoprofile**
The tumour is diffusely immunoreactive for S100 protein, but the proportion of positive cells is lower than that in schwannoma. CD34 stains the admixed fibroblasts.

**Genetic susceptibility**
Sinonasal neurofibromas are generally not associated with NF1 [1091,1095, 2018].

**Prognosis and predictive factors**
Neurofibromas are benign and have a very low recurrence rate. A small percentage of cases may undergo malignant transformation.

**Meningioma**

**Definition**
A benign neoplasm of meningothelial cells.

**ICD-O code**

**Epidemiology**
Primary extracranial (ectopic, extracalvarial) meningiomas of the sinonasal tract are rare, comprising <0.5% of non-
Tumours of the nasal cavity and paranasal sinuses

epithelial neoplasms {814,873,1109,1781,2019,2221,2599}. They should be distinguished from intracranial meningiomas with extracranial/extraspinal extension into the sinonasal tract {721, 814,2599}. Any age can be affected, and there is a slight female predilection. Men tend to be younger than women by about a decade.

Localization
Sinonasal tract meningiomas involve both the nasal cavity and paranasal sinuses more frequently than either location alone. Most tumours are left-sided {814,873,1109,1781,2019,2221,2599}.

Clinical features
Symptoms include a mass (often polypoid), nasal obstruction, epistaxis, sinusitis, pain, headache, seizure, exophthalmos, periorbital edema, visual disturbance, ptosis, and facial deformity {814,873,1109,1781,2019,2221,2599}. Symptoms are present for an average of 4 years.

Macroscopy
The tumours range up to 8 cm, with a mean of about 3 cm. They may infiltrate bone and rarely ulcerate the mucosa. The cut surface is grey-white, tan or pink, gritty, firm to rubbery. Calcifications and fragments of bone are frequently visible.

Tumour spread and staging
Primary extra-cranial meningiomas have not been reported to metastasize {814, 873,2019,2599}.

Histopathology
Sinonasal meningiomas can exhibit a variety of histological patterns, most commonly meningotheliomatous, characterized by lobules of cells with whorl formation, indistinct cell borders, and bland nuclei with delicate chromatin {1329,2599}. Intranuclear pseudoinclusions and psammoma bodies are common. Other variants can also occur in the sinonasal tract, such as transitional, metaplastic (lipidized cells within tumour), and psammomatous type {1329}.

Immunoprofile
Meningiomas are immunoreactive for epithelial membrane antigen and vimentin, but usually negative for cytokeratin, although rare lesions can exhibit focal and weak cytokeratin immunoreactivity. They are frequently positive for progesterone receptor (50%) and occasionally for oestrogen receptor (25%). Glial fibrillary acidic protein and smooth muscle actin are negative.

Differential diagnosis
The differential diagnoses include carcinoma, melanoma, aggressive psammomatoid ossifying fibroma and follicular dendritic cell sarcoma/tumour {2599, 2771}.

Histogenesis
Meningiomas are derived from arachnoid cap cells located extra-cranially within the sheaths of nerves or vessels.

Prognosis and predictive factors
Complete surgical extirpation is sometimes difficult, and accounts for the up to 30% recurrence rate {1109,2019,2599}. The rare deaths are related to compromise of mid-facial structures or complications of surgery, rather than the aggressive nature of the tumour. Histologic features (such as hypercellularity, nuclear pleomorphism, necrosis), proliferation index and progesterone receptor status do not influence prognosis {1138,1139,1426,1666,2599}. 
Chondrosarcoma, including mesenchymal chondrosarcoma

Definition
Chondrosarcoma is a malignant tumour of hyaline cartilage. Mesenchymal chondrosarcoma is a malignant small round cell neoplasm with focal cartilaginous differentiation, and often with a pericytomatosus vascular pattern.

ICD-O codes
Chondrosarcoma 9220/3
Mesenchymal chondrosarcoma 9240/3

Synonym
Polyhistioma

Epidemiology
These tumours are rare in the facial skeleton. Chondrosarcomas account for <16% of all sarcomas of the nasal cavity, paranasal sinuses and nasopharynx (256,463,1367,4045). Chondrosarcoma affects older adults, with a male predilection. Mesenchymal chondrosarcoma is extremely rare, and affects young adults, with a female predilection.

Localization
Chondrosarcoma involves the alveolar portion of the maxilla, the maxillary sinus or the nasal septum. Mesenchymal chondrosarcoma involves the mandible and maxilla almost equally.

Clinical features
Patients with involvement of the nose present with nasal obstruction. Painful swelling is common with other sites of involvement.

Imaging
On plain radiographs, both tumours show osteolysis with stippled calcification, cortical destruction and possible soft tissue extension. Computerized tomograms and magnetic resonance images are useful in evaluating the extent of disease (463).

Macroscopy
Chondrosarcomas are lobulated pale-blue glistening masses that may show cystic change. Mesenchymal chondrosarcomas have the fish-flesh appearance of high-grade sarcomas; chalky foci of calcification may offer a diagnostic clue.

Histopathology
Chondrosarcomas are often lobulated, and show round to oval cells in lacunae with a blue chondroid matrix that may show myxoid changes. Most are low-grade. Increased cellularity and permeation of the inter trabecular spaces of bone, if identified, are the most important features that distinguish chondrosarcoma from chondroma. Radiological correlation is required for a definitive diagnosis (4045). Mesenchymal chondrosarcomas show a mixture of hyaline cartilage and small round to oval cells with hyperchromatic nuclei, frequently arranged in a pericytomatosus vascular pattern. These cells...
Tumours of the nasal cavity and paranasal sinuses

are frequently immunoreactive for CD99. The relative amounts of the two elements are quite variable. The chondroid lobules have the appearance of well-differentiated chondrosarcoma.

**Prognosis and predictive factors**

Chondrosarcomas are associated with an excellent prognosis if the lesions are completely resected. Approximately 20% of patients die of tumour, most often with uncontrolled local recurrence (2223,4045).

Mesenchymal chondrosarcoma is a high-grade tumour with an unpredictable prognosis. Patients with tumour of the facial skeleton do better than those with tumours of the remainder of the skeleton (2687).

**Osteosarcoma**

**Definition**

Osteosarcoma is a primary malignant tumour of bone in which the neoplastic cells produce osteoid or bone.

**ICD-O code** 9180/3

**Synonym** Osteogenic sarcoma

**Epidemiology**

Osteosarcomas of the jaws are very rare, with an incidence of 0.7 per million (868). They are extremely rare in other head and neck sites. Patients are a decade older than those with extragnathic osteosarcomas (455,868,1366). There is a modest male predilection.

**Etiology**

Over 10% of tumours are post-radiation, including Thorotrast exposure.

**Localization**

The maxilla and the mandible are affected almost equally. In the maxilla, the alveolar ridge and the antrum are predominantly involved, whereas in the mandible, the body is the main site.

**Clinical features**

Symptoms include swelling with or without pain and loosening of teeth.

On plain radiograph, the tumour is usually lytic but may be sclerotic or mixed. In over half of the lesions, there is soft tissue extension. Computerized tomogram is better in demonstrating matrix mineralization and soft tissue extension (1457).

**Macroscopy**

The tumours vary from the lobulated blue colour of cartilage to fleshy white to densely sclerotic masses.

**Histopathology**

Osteosarcomas of the jaws are generally better differentiated than extragnathic osteosarcomas. There is commonly chondroblastic differentiation, characterized by lobules of atypical-appearing chondrocytes in lacunae. There is a typical condensation of nuclei toward the periphery of the lobules, where sheets of spindle cells may be seen. The centre of the chondroid lobules shows bone formation in the form of trabeculae. The remainder show osteoblastic or fibroblastic features. It is unusual to see benign giant cells within the tumour.

**Prognosis and predictive factors**

Some studies have shown that patients with osteosarcoma of the jaws have a better survival than those with extragnathic osteosarcomas (455,1366). However, some other studies (206,868) have not confirmed this finding. Complete surgical resection is associated with better prognosis.
Benign tumours of bone and cartilage

**Fibrous dysplasia**

See Chapter 6 for details.

**Giant cell lesion**

**Synonym**

Giant cell granuloma.

Extragnathic giant cell lesion may rarely involve the paranasal sinuses [2161, 2243,2479,2648,2839]. Symptoms include pain, visual disturbances, exophthalmos, epistaxis, lacrimation, anaesthesia and swellings [728,1867,2322,2839]. Plain radiographs show nonspecific osteolytic defects. Their extent and possible impingement on brain or orbital contents is better visualized in cross-sectional studies (CT, MRI) [2322,2839]. Brown tumour of hyperparathyrodism should be excluded. Please refer to Chapter 6 for details.

**Giant cell tumour of bone**

**Definition**

An aggressive but benign neoplasm containing spindle-shaped stromal cells, mononuclear round to oval cells resembling histiocytes, and abundant evenly distributed osteoclastic giant cells.

**ICD-O code**

9250/1

**Epidemiology**

Giant cell tumour (GCT) accounts for about 5% of all bone tumours. Most cases occur in patients between 20 and 50 years [523,775,1146,1369]. CGT is slightly more common in women than in men [523,775,2831].

**Localization**

In the skull, in the absence of Paget disease, the bones developing from endochondral ossification, i.e. sphenoid, ethmoid and temporal bone, are almost exclusively involved [207,496,870,875].

**Clinical features**

**Signs and symptoms**

Clinical symptoms of GCTs depend on the site of occurrence. Sphenoidal lesions are associated with headache, diplopia and vision impairment or cranial nerve palsies (II, III, IV, V, VI and combinations). Temporal bone involvement causes deafness (conductive; middle ear and mastoid; sensorineural combined with vertigo: petrous bone), retroauricular pain, or swelling [676,2328]. Duration of symptoms ranges from weeks to years [207].

**Imaging**

On plain radiographs and CT, GCT presents with a nonspecific expansile and destructive osteolysis, generally lacking any matrix mineralization. The lesion shows contrast enhancement on CT [207,2177,2806]. A soft tissue mass in the sphenoid sinus or the sella turcica, displacing the pituitary gland, is best seen on cross-sectional studies (CT, MRI) [983,2177,2556].

**Histopathology**

The tumour is characterized by abundant multinucleated osteoclastic giant cells, with up to 50-100 nuclei, that are evenly distributed among sheets of stromal cells. In some areas, ovoid to plump spindled stromal cells are more prominent, and giant cells may be lacking. Regressive changes, including fibrosis, foam cell aggregates, haemosiderin deposits, and even necrosis may be present. Small foci of reactive woven bone are often seen. Mitoses are easily

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Fig. 1.67 Giant cell tumour. Large geographic osteolytic lesion (OL) at the base of the skull with an irregular dorsal margin.

Fig. 1.68 Central giant cell lesion/granuloma. A Osteodestructive lesion of the right maxillary sinus extending into the nasal cavity, eroding the orbital floor and destroying the alveolar process of the maxilla with protrusion into the oral cavity. B On T1 images the signal intensity is low. C Fluid levels are indicative for a pseudocystic (ABC-like) component.
found in the mononuclear cells, but atypical ones do not occur and, if present, are a strong indicator for progression to malignant GCT [272]. Intravascular growth, particularly in the tumour periphery, may be noted, but has no prognostic relevance [207,775,2831].

Histogenesis
The ovoid to plump spindled stromal cells represent the active proliferating tumour cell pool [2190], capable of secreting cytokines and differentiation factors, including receptor activator of nuclear factor κ ligand (RANKL) [2207]. These factors attract monocytes, the second cell type in GCT, and promote fusion to osteoclasts, the third cell type in GCT. The monocytes and osteoclast-like giant cells represent a non-neoplastic tumour component [775,2831].

Genetics
Cytogenetic studies reveal telomeric associations (TAS) as the most frequent chromosomal aberration [272]. Some GCTs show rearrangements in 16q22 or 17p13, similar to aneurysmal bone cyst, which is often associated with GCT. These rearrangements may indicate the possible presence of an aneurysmal bone cyst component [775].

Prognosis and predictive factors
GCT of the skull is a locally aggressive lesion. Treatment consists of complete removal, if possible. Radiotherapy is also applied [207,1146]. Histologically malignant GCT of the skull has been rarely described, sometimes associated with Paget disease [365,1474].

**Chondroma**
Chondromas of the sinonasal tract are extremely rare, and any cartilaginous tumour greater than 2 cm occurring in this site should be considered potentially malignant until proven otherwise.

**ICD-O code** 9220/0

**Osteoma**

**Definition**
A benign lesion composed of mature bone with a predominantly lamellar structure.

**ICD-O code** 9180/0

**Synonyms**
In the jaws and calvaria, the terms exostosis and osteoma have been used interchangeably [131]. The term ‘osteoma’ should be used in a restricted sense limited to lesions of the paranasal sinuses, facial bones and orbit, although it has been used in the literature to describe calvarian and mandibular ivory exostosis, surface (juxtapositional) osteoma of the long bones, torus palatinus and torus mandibularis.

**Epidemiology**
Among patients with sinonasal radiographs taken for a variety of reasons, up to 1% have been found to have osteomas. It may occur at any age, but especially in young adults. There is a 2:1 male predominance.

**Localization**
The osteomas may be single or multiple; central or on the bone surface, where they can be sessile or rarely pedunculated. They occur most commonly in the frontal and ethmoid sinuses. The maxillary and sphenoid sinuses are infrequently involved. In the jaws, the angle of the mandible is more frequently involved than the coronoid process or condyle.

**Clinical features**
Osteomas are often asymptomatic and incidentally discovered. However they can produce pain or symptoms related to the location. Multiple jaw osteomas are a frequent component of the Gardner syndrome (a form of familial adenomatous polyposis), being found in 70-90% of patients. Osteomas are radiodense, sharply defined, well-circumscribed lesions occurring in either a central or peripheral location.

**Macroscopy**
The lesion is a well-circumscribed white bony mass, which is occasionally polypoid or exophytic.

**Histopathology**
Osteoma is characterised by compact cortical bone with scanty intervening fibrovascular stroma. In some cases, there is a peripheral rim of dense sclerotic lamellar bone surrounding trabeculae of lamellar or occasionally woven bone separated by fibrofatty vascular tissue.

**Genetic predisposition**
The presence of multiple osteomas is an
important clue that the patient may have Gardner syndrome.

**Prognosis and predictive factors**
No therapy is required unless the lesion causes cosmetic or functional problems. A local resection is the treatment of choice in such circumstances.

**Chondroblastoma and chondromyxoid fibroma**

**ICD-O codes**
- Chondroblastoma 9230/0
- Chondromyxoid fibroma 9241/0

Chondroblastomas and chondromyxoid fibromas are rare in the head and neck (177,988,1024,1120,1185,1348,1349,1356,1466,2559,2683,2730). See WHO Classification of Tumours of Soft Tissue and Bone [775].

**Osteochondroma (exostosis)**

**Definition**
A pedunculated or sessile exophytic bony projection with a cartilaginous cap. The bony component is continuous with the underlying bone.

**ICD-O code**
9210/0

**Epidemiology**
Osteochondroma is one of the most common lesions of the long and flat bones. It may be solitary or multiple. In the facial bones, osteochondromas are very rare and almost invariably single. Osteochondromas of the facial skeleton have not been reported in the setting of multiple hereditary exostoses. The mean age at diagnosis is 40 years, which is older than that of patients with tumours occurring outside the head and neck. Females are more commonly affected than males.

**Localization**
More than half of the lesions occur in the coronoid process of the mandible. The condyle can also be involved.

**Clinical features**
Osteochondroma involving the coronoid process or the condyle causes difficulty in opening the mouth or dysfunction of the temporomandibular joint. On plain radiographs, flaring of the cortex in continuity with the underlying bone and varying degrees of calcification and/or ossification are present. The cartilaginous cap is of variable thickness.

**Macroscopy**
A cartilaginous cap covers the bony protrusion.

**Histopathology**
The cap consists of hyaline cartilage, and the osteochondral junction resembles the growth plate. A well-defined zone of enchondral ossification matures into cancellous bone with marrow.

**Prognosis and predictive factors**
Excision is curative. No recurrence or malignant transformation has been reported in osteochondromas of the jaws.

**Osteoid osteoma**

**ICD-O code**
9191/0

**Definition**
A rare, benign bone-forming tumour of limited growth potential, usually less than 1.5 cm, typically associated with nocturnal pain that is relieved by salicylates. It is very rare in the head and neck. It occurs in young patients (first three decades), with male predominance. On plain radiographs, dense cortical sclerosis surrounds a radiolucent nidus. Histologically, the nidus shows interconnected, ossified woven bone rimmed by osteoblasts. Fibrous tissue, vessels and multinucleated giant cells are identified in between the bony trabeculae. See WHO Classification of Tumours of Soft Tissue and Bone [775].

**Osteoblastoma**

**ICD-O code**
9200/0

**Definition**
A rare, benign, bone-forming tumour in which osteoblasts rim woven bony trabeculae, forming a mass usually over 2 cm.

**Epidemiology**
Osteoblastoma is rare, and 90% of cases occur below the age of 30 years. It is more common in males.

**Localization**
In the head and neck, the most common site of involvement is the jaws, followed by the cervical vertebrae and the skull (1570). The mandible is affected about two to three times more often than the maxilla. Most arise in the body of the mandible, rarely in the midline or coronoid process.

**Clinical features**
Osteoblastomas of the jaw cause swelling and toothache, and in the cervical spine, pain, scoliosis and nerve root compression. In contrast to osteoid osteoma, the pain is rarely nocturnal and not relieved by salicylates.

---

**Fig. 1.70**

A Osteoma of the right parietal bone. Mature lamellar bone with osteon-like structures. B Osteoid osteoma. Osteoblasts surround the trabeculae. C Nidus of an osteoblastoma, showing a single layer of osteoblasts lining the bony trabeculae.
On plain radiographs, osteoblastoma is a sharply circumscribed, oval-round lytic lesion. It may have a mixed lytic and sclerotic pattern, reflecting the degree of mineralization of the matrix. A reactive bony shell is detected at the periphery. Radiographic features indistinguishable from malignant lesions are reported in about one-third of cases.

**Macroscopy**
It is a red and gritty lesion often with cyst formation. The border between the tumour and the host bone is very sharp.

**Histopathology**
Woven bony trabeculae, rimmed by osteoblasts are haphazardly distributed within a richly vascularized fibrous stroma accompanied by osteoclast-like giant cells. Mitotic figures may be present, but without atypical forms. Degenerative nuclear atypia is occasionally present. When large plump osteoblasts with prominent nuclei predominate, the tumour is often referred to as epithelioid osteoblastoma or aggressive osteoblastoma. However, these histologic features are not necessarily indicative of aggressive behaviour. Rarely, focal areas of hyaline cartilage may be identified, as well as secondary aneurysmal bone cyst-like changes. At the periphery, there is no permeative growth pattern. The histologic features are identical to cementoblastoma. Tumours showing direct continuity with the root of a tooth are preferably termed a cementoblastoma.

**Prognosis and predictive factors**
Curettage or local excision is the treatment of choice. In the few cases with recurrence, a further conservative treatment will control the disease.

**Ameloblastoma**
Ameloblastomas are very rare in the sinonasal tract and nasopharynx (1554, 2257). See Chapter 6 on odontogenic tumours for details.

**ICD-O code** 9310/0

**Nasal chondromesenchymal hamartoma**

**Definition**
A tumefactive process arising in the nasal cavity and/or paranasal sinuses whose mixed chondroid, stromal, and cystic features are morphologically similar to the chest wall hamartoma.

**Epidemiology**
There are only 12 reported cases (45,1140,1284,1311,1678). A pleuropulmonary blastoma was diagnosed in one of these children (1678). One infant with the prenatal detection of hydrocephalus also had absence of the corpus callosum and hypoplasia of the cerebellar vermis. The age range is newborn to 16 years with most cases presenting in the first year of life, often before 3 months of age. There is a male predilection of approximately 3:1.

**Clinical features**

**Signs and symptoms**
Respiratory difficulty, the discovery of an intranasal mass and/or facial swelling are the most common presenting features. The respiratory distress is detected in the immediate neonatal period or develops later during feedings with accompanying cyanosis. A unilateral mass in the nasal cavity is the most consistent finding on physical examination.

**Imaging**
A mass density in the nasal cavity and/or the contiguous paranasal sinuses is noted on radiographic examination. Magnetic resonance or computed tomographic imaging discloses a dense mass with or without calcifications or a heterogeneous signal in a lesion with cystic features. Extension or involvement of the maxillary and/or ethmoid sinuses and erosion into the anterior cranial fossa are other accompanying changes.

**Macroscopy**
Multiple solid and cystic fragments of tis-
sues, some with identifiable foci of cartilage reflect the piecemeal nature of the resection in most cases. The precise site of origin of the mass has varied from the nasal septum, upper nasal cavity or floor of the anterior cranial fossa.

**Histopathology**
All tumours have had nodules of cartilage varying in size, contour and degree of differentiation. Some nodules resemble the chondromyxomatous nodules of a chondromyxoid fibroma, whereas others are well-differentiated cartilaginous nodules. At the periphery of the chondroid nodules, there is a loose spindle cell stroma or an abrupt transition to hypocellular fibrous stroma. Other areas can have a fibro-osseous appearance with a prominent cellular stromal component and small ossicles or trabeculae of immature woven bone resembling fibrous dysplasia. Yet another common pattern is a cellular stroma with hyalinized nodules with or without perivascular stromal cells displaying a pericytomatosous pattern. Cellular myxoid foci are similar in some respects to cranial/nodular fasciitis. The aneurysmal bone cyst-like areas are surrounded by a stroma rich in multinucleated giant cells.

**Immunoprofile**
The cartilage, mature or immature, is immunoreactive for S-100 protein. The spindled stroma is immunoreactive for smooth muscle actin and vimentin.

**Differential diagnosis**
The differential diagnosis depends on the particular combination of microscopic features present in the biopsy or resection. Since cartilage is the dominant component, differential diagnoses include chondromyxoid fibroma and chondroblastoma (1858). Other differential diagnoses may include aneurysmal bone cyst, fibrous dysplasia, cranial fasciitis and osteochondromyxoma (334). Interestingly, the latter tumour may be congenital, may involve the paranasal sinuses and is associated with the Carney complex in some cases. None of the patients with nasal chondromesenchymal hamartoma are known to have the Carney complex (1678).

**Prognosis and predictive factors**
Information on the clinical behaviour is incomplete, but prognosis is apparently favourable. There is some capacity for continued local growth when the resection is incomplete.
Tumours of the nasal cavity and paranasal sinuses

A.C.L. Chan
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S.B. Kapadia

Haematolymphoid tumours

Non-Hodgkin lymphoma

Definition
Primary non-Hodgkin lymphomas (NHL) of the nasal cavity or paranasal sinuses are defined as lymphoid cell neoplasms in which the bulk of disease occurs in these anatomic sites.

Synonyms
Most cases described in the past as polymorphic reticulosis, malignant midline reticulosis, lethal midline granuloma or angiocentric immunoproliferative lesion, are now reclassifiable as extranodal NK/T cell lymphoma of nasal-type.

Epidemiology
Malignant lymphoma is the second most common malignancy of the nasal cavity and paranasal sinuses, following squamous cell carcinoma [1013]. It accounts for 14% of all cancers in these sites [1013]. Although many different types of NHL can occur in the nasal cavity, the most common lymphoma type is extranodal NK/T cell lymphoma of nasal-type, especially in Asian populations [5,420]. The relatively high prevalence of this lymphoma type in Asians and Latin Americans also accounts for the higher overall incidence of nasal lymphomas in these populations as compared with Caucasian populations [58,2104]. Other peripheral T-cell lymphomas, such as anaplastic large cell lymphoma, can also occur in the sinonasal region.

Lymphomas presenting in the paranasal sinuses are frequently B-cell lymphomas, with diffuse large B-cell lymphoma (DLBCL) being the most common [5,551,1837]. Other B-cell lymphomas that can involve the sinonasal regions include Burkitt lymphoma, follicular lymphoma, extranodal marginal zone B-cell lymphoma of MALT type, and mantle cell lymphoma [5]. Please also refer to the section of non-Hodgkin lymphomas in Chapter 4 on tumours of the oral cavity and oropharynx.

NHL of the nasal cavity and paranasal sinus is primarily a disease of adults, with male predominance. Patients with extranodal NK/T cell lymphoma of nasal-type have a male to female ratio of 3:1, and a median age of 53 years [420]. Patients with DLBCL are generally one decade older (median age 63 years), and the male to female ratio is 1.2:1 [5,420]. Children may rarely present with NHL of the nasal cavity and the paranasal sinuses, with Burkitt lymphoma being the most common type [2808].

Etiology
The etiology is unknown, but extranodal NK/T cell lymphoma of nasal-type is strongly associated with Epstein-Barr virus (EBV) irrespective of the ethnic background of the patients [68,376,1107,1347,2671,2828]. There is only a weak association between B-cell lymphomas in the nasal cavity and the paranasal sinuses with EBV [376,511,2617,2740]. Immunosuppression (e.g. post-transplant, HIV infection) is associated with an increased risk of developing NHL, including in the nasal cavity and paranasal sinuses. Although the majority of the cases in immunosuppressed patients are DLBCL [511,2068], extranodal NK/T cell lymphoma of nasal-type has also been reported [328]. Most of the NHL that arise in the setting of immuno-suppression are also EBV-related [511].

Localization
Lymphomas of the nasal cavity are often locally destructive, with obliteration of the nasal passages and maxillary sinuses. In particular, extranodal NK/T cell lymphoma can involve the adjacent alveolar bone, hard palate, orbit and nasopharynx in over half of the cases [1948]. Lymphomas of the paranasal sinuses commonly show bony destruction and local extension to adjacent structures including the orbit, palate, nasal cavity, nasopharynx, and soft tissues in the cheek and infratemporal fossa [511,1836,1837]. Maxillary sinus is the most commonly involved paranasal sinus.

Clinical features
Patients may present with nasal obstruction, epistaxis, nasal discharge, pain and nasal swelling or facial swelling. Locally advanced cases can cause destruction of midline facial structures. The nasal septum or palate may be perforated. Extension to the orbits can lead to proptosis and visual disturbance. Regional lymph node involvement may occur in some patients. Occasional patients have systemic symptoms including fever and weight loss. Haemophagocytic syndrome with pancytopenia occurs at presentation in a minority of patients with extranodal NK/T cell lymphoma of nasal-type [420,2533].
Haematolymphoid tumours

**Tumour spread and staging**

The majority (80%) of patients with extranodal NK/T cell lymphoma of nasal-type have localized disease at presentation (Stage IE/IIE) \[420,1500,1505\]. Bone marrow involvement at presentation is uncommon \[2810\]. Although extranodal NK/T cell lymphoma often shows localized disease at presentation, spread to other sites (such as skin, gastrointestinal tract, liver, lymph node, testis) during the course of disease is common. Most of the patients (75%) with DLBCL of the nasal cavity and the paranasal sinuses present with low clinical stage (IE/IIE) \[420,511\]. In contrast to extranodal NK/T cell lymphoma, cervical lymph node involvement is more frequent at presentation (60%), and the common sites of relapse are lymph node, liver and lung \[420\].

**Extranodal NK/T cell lymphoma**

**ICD-O code** 9719/3

Extranodal NK/T cell lymphoma of nasal-type is characterized by a diffuse lymphomatous infiltrate expanding the nasal or paranasal sinus mucosa, with wide separation and destruction of the mucosal glands, which may undergo a peculiar clear cell change. Extensive coagulative necrosis and frequent apoptotic bodies are very common, as are ulceration, angiocentricity, angiodestruction and fibrinoid deposits in vessel walls. The lymphoma cells vary in size in different cases, ranging from small through medium-sized to large. Some nuclei have an irregular outline, while others can be round or oval. The cells have a moderate amount of pale cytoplasm, and cytoplasmic azurophilic granules can be identified in Giemsa-stained touch preparations. Some cases are associated with a rich inflammatory infiltrate, consisting of small lymphocytes, histiocytes, plasma cells and eosinophils. Occasionally pseudoepitheliomatous hyperplasia of the overlying squamous epithelium may occur, mimicking well differentiated squamous cell carcinoma.

**Immunoprofile and genetics**

The lymphoma most commonly exhibits an NK-cell immunophenotype of CD2+, surface CD3(Leu4)-, cytoplasmic CD3+, CD56+ \[1196\]. CD43 and CD45RO are commonly positive, but other T-cell markers (including CD5) and NK-cell markers (CD16, CD57) are usually negative. The tumours commonly exhibit a cytotoxic phenotype with expression of perforin, TIA1, and granzyme B \[662,1773,1878,1935\]. Fas (CD95) and Fas ligand expression are frequent, and may account for the extensive necrosis \[1877,1935\]. Expression of the various NK cell receptors is variable. P-glycoprotein/MDR1 is often expressed (90%), and may explain the poor response to chemotherapy \[2838\]. The T-cell receptor genes are often in germline configuration \[1196\]. Practically all cases (>95%) are associated with EBV \[68,376,1107,1263,1347,2671,2828\]. The virus is best demonstrated in the tumour cells by in situ hybridization for EBER (EBV-encoded early RNAs) \[376\]. The EBV is in clonal episomal form, pro-

---

**Fig. 1.76** Nasal NK/T cell lymphoma. **A** In this example, the mucosa is intact and expanded by a diffuse infiltrate of lymphoma cells. **B** The mucosal lymphoid infiltrate is destructive, resulting in separation and loss of mucosal glands.

**Fig. 1.77** Nasal NK/T cell lymphoma. **A** This case shows marked pseudoepitheliomatous hyperplasia, mimicking squamous cell carcinoma. **B** Note angiocentric and angiodestructive growth.
viding additional evidence of the clonal nature of the lesion \(\{1107,1693\}\). Some cases are CD56 negative, but are still classified as extranodal NK/T cell lymphoma provided they express T-cell markers and cytotoxic markers, and are EBV positive. These cases may show clonally rearranged T-cell receptor genes and may represent a neoplasm of cytotoxic T-lymphocytes \(\{426\}\). T-cell lymphomas which lack cytotoxic markers or evidence of EBV infection are diagnosed as peripheral T-cell lymphoma unspeciﬁed. Lymphoblastic lymphoma of probable NK-cell lineage (or so-called blastic NK-cell lymphoma) with expression of CD56 and TdT and no EBV association has also been described in the nasal cavity, but this is an entity distinct from the extranodal NK/T cell lymphoma of nasal-type \(\{1352,1838\}\).

**Differential diagnosis**

Since the tumour cells of extranodal NK/T cell lymphoma can be masked by a prominent inﬁltrative lymphoid pattern, the lesion can be mistaken as an infective, inﬁltrative or granulomatous lesion (including Wegener granulomatosis). It is not uncommon that a deﬁnitive diagnosis can only be reached after repeated biopsies. While Wegener granulomatosis similarly presents with destructive nasal lesion, simultaneous pulmonary involvement may be present. There is serologic positivity for cytoplasmic anti-neutrophil cytoplasmic antibody (c-ANCA), and the main histologic ﬁndings are chronic inﬁammation with microabscesses and histiocytic inﬁltrate, in the absence of atypical lymphoid cells.

In those examples of extranodal NK/T cell lymphomas dominated by small cells with minimal atypia, a deﬁnitive diagnosis can be difﬁcult to make. An angiocentric inﬁltrate with expansion of the mucosa and mucosal gland destruction, coupled with prominent necrosis, should raise suspicion for the diagnosis of extranodal NK/T cell lymphoma. Conﬁrmation of the diagnosis can be made by demonstrating sheets of CD3+, CD56+ and EBER+ cells. Some non-lymphoid CD56+ small round cell tumours (e.g. olfactory neuroblastoma, Ewing sarcoma/primitive neuroectodermal tumour, rhabdomyosarcoma) also enter in the differential diagnoses. However, these can be easily excluded by appropriate immunohistochemical stains.

**Diffuse large B-cell lymphoma**

ICD-O code 9680/3

In DLBCL of the nasal cavity or paranasal sinuses, the mucosa shows dense, diffuse and interstitial inﬁltration by large or medium-sized lymphoid cells. There may or may not be ulceration and necrosis. Occasional cases show angioinvasion. The tumour cells may resemble centroblasts or immunoblasts, or have a non-speciﬁc blastoid appearance. The nuclei are round, multilobated or irregularly folded, with multiple small membrane-bound nucleoli or single central prominent nucleolus. The tumour cells express pan-B markers (e.g. CD20, CD79a). Extramedullary myeloid sarcoma, plasmacytoma, undifferentiated carcinoma and amelanotic melanoma may resemble DLBCL, but these entities can be readily distinguished by appropriate immunohistochemical stains.

**Histogenesis**

Most cases of extranodal NK/T cell lymphoma of nasal-type are activated NK-cell neoplasms, while some appear to be neoplasms of cytotoxic T-cells \(\{425\}\). DLBCL are mature B-cell neoplasms at either the germinal centre or post-germi nal centre stage of differentiation.

**Somatic genetics**

A number of cyogenetic abnormalities

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Table 1.4  Non-Hodgkin lymphomas in the nasal cavity or paranasal sinuses: differences in distribution according to cell lineage.

<table>
<thead>
<tr>
<th></th>
<th>Primary in nasal cavity</th>
<th>Primary in paranasal sinuses</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>NK/T- or T-cell lymphomas</td>
<td>B-cell lymphomas (mostly DLBCL)</td>
</tr>
<tr>
<td>Asian series ({420,1837})</td>
<td>71%</td>
<td>29%</td>
</tr>
<tr>
<td>Western series ({5})</td>
<td>54%</td>
<td>46%</td>
</tr>
</tbody>
</table>

DLBCL = diffuse large B-cell lymphoma
Haematolymphoid tumours

have been reported in extranodal NK/T cell lymphoma of nasal-type, most commonly isochromosome 1q, isochromosome 6p, partial deletion of 6q, and aberration at 11q [2606,2811]. Comparative genomic hybridization and loss of heterozygosity studies have suggested frequent DNA loss at 1p, 6q, 11q, 12q, 13q, 17p, whole X, and frequent gain at 1p, 2q, 6p, 10q, 11q, 12q, 13q, 17q, 19p, 20q, Xp [1346,2382,2383]. Overall, the most frequent changes are del(6)(q21-25), del(17)(p12-p13), del(13)(q14-q34) and gain of 1p32-pter [422]. P53 protein overexpression occurs in 45-86% of cases [1496,2104,2105], but P53 mutation is less common (24-48%) [1496,2105]. TP53 mutation has been associated with large cell morphology and advanced stage [2105]. FAS gene mutation is frequently observed [2331,2534]. Aberrant methylation of promoter CpG region of P73 gene occurs in 94% of cases [2381], and its detection by methylation-specific polymerase chain reaction may be helpful for monitoring residual disease or early relapse [2380]. There are no molecular data on DLBCL specifically occurring in the sinonasal tract.

Genetic susceptibility
Extranodal NK/T cell lymphoma of nasal-type has been reported in both the father and son of a family with known pesticide exposure [1354].

Prognosis and predictive factors
Radiotherapy and/or systemic chemotherapy is the treatment of choice for localized disease [420-422,1505,1550]. Treatment of DLBCL follow protocols for similar tumours elsewhere in the body, as some series showed that chemotherapy might be beneficial [1550,2091]. The overall survival for extranodal NK/T cell lymphoma of nasal-type is only 30-50% [420-422,1312,1838]. In patients achieving complete remission, local relapse occurs in one-third to one-half of cases [421,1312], and systemic failure is also common [421]. Factors associated with a worse outcome include: advanced stage, poor performance status, B symptoms, and bulky disease [422]. There is no conclusive evidence to suggest that the histologic grading of NK/T cell lymphoma can predict the clinical outcome. Expression of cutaneous lymphocyte antigen (CLA) may be associated with a worse prognosis, but this finding has yet to be confirmed [2863]. The prognosis is slightly more favourable for DLBCL compared with extranodal NK/T cell lymphoma of nasal-type [420]. The overall survival for DLBCL is 35-60% [420,511,1550]. Prognostic factors have not been studied in detail in sinonasal DLBCL. A Western series reporting treatment results of lymphomas of the nasal cavity and the paranasal sinuses showed that the International Prognostic Index is the only significant predictor for freedom from progression rate [1550].

Extramedullary plasmacytoma

Definition
A mass-forming lesion of monoclonal plasma cells that occurs outside the bone and bone marrow. By definition, patients with primary extramedullary plasmacytoma (EMP) do not have evidence of underlying multiple myeloma.

ICD-O code 9734/3

Epidemiology
The mean age of patients with EMP of the head and neck is 60 years (range 34-78 years), with a male predominance 4:1 [1267].

Localization
Most frequent sites of involvement are nasal cavity, paranasal sinuses and nasopharynx [433,827,1267,1613,1972,2347,2656,2746].
Clinical features
EMP tends to be solitary, with multiple tumours present in only 10% of cases at diagnosis. The presenting features of head and neck EMP are: soft tissue mass (80%), airway obstruction (35%), epistaxis (15%), nasal discharge (10%), regional lymphadenopathy (10%), and cranial nerve palsies (5%). The mean duration of symptoms is about 4.5 months. The tumour ranges in size from 2-5 cm. The appearance varies from grey to red, soft to firm, and sessile or pedunculated. EMP bleeds easily and is usually smooth without mucosal ulceration. Cervical lymph nodes are enlarged in only 10% of patients (2500). Occasional primary EMP may be associated with serum paraproteinaemia. An underlying multiple myeloma should always be excluded.

Macroscopy
EMP is lobulated, smooth or nodular, and has a fleshy or rubbery consistency.

Histopathology
There is a diffuse infiltrate of neoplastic plasma cells in the subepithelial tissue, accompanied by a scant vascularized stroma, and rarely blood lakes. There can be deposits of amyloid or immunoglobulin in the stroma. The tumour can be well, moderately or poorly differentiated [18,1267]. Well-differentiated EMP is characterized by uniform normal-looking to mildly atypical plasma cells. Intracytoplasmic crystals can be abundant in some cases. Dutcher bodies are sometimes seen. Moderately-differentiated EMP comprises moderately atypical plasma cells that vary in size. Poorly-differentiated (anaplastic) EMP comprises large cells that are often barely recognizable as being plasma cells. The nuclei often show significant variation in size, and can be round or irregularly folded. The chromatin pattern ranges from vesicular to finely granular to coarsely clumped. Nucleoli can be prominent. The cytoplasm is amphophilic and eccentrically located, and a paranuclear hof (Golgi zone) may be present. Mitotic figures are frequent. Some tumour cells can be multinucleated.

Differential diagnosis
Well-differentiated EMP should be distinguished from reactive plasma cell proliferations, either non-specific or associated with specific disorders, such as rhinoscleroma or Rosai-Dorfman disease. Reactive plasmacytic proliferations show a polyclonal pattern of immunoglobulin staining. Moderately or poorly-differentiated EMP can cause significant difficulties in distinction from large cell lymphoma, carcinoma, melanoma, extramedullary myeloid sarcoma and olfactory neuroblastoma. The occasional positive staining for cytokeratin can lead to a misdiagnosis of carcinoma. A high index of suspicion for EMP should be raised for any poorly differentiated neoplasm occurring in the upper aerodigestive tract. Features suggestive of the diagnosis include eccentrically placed nuclei, coarsely clumped “clock-face” chromatin in some nuclei, and amorphophilic cytoplasm with a paranuclear hof. The diagnosis can be confirmed by immunohistochemistry or in-situ hybridization for immunoglobulin mRNA to look for monotypic light chain expression [18].

Prognosis and predictive factors
The mainstay of treatment for primary EMP is radiotherapy. The prognosis of primary EMP is far more favourable than that associated with myeloma [1267]. Approximately 20% of patients with primary EMP will develop multiple myeloma, but it is not possible to predict which cases will progress.

Extramedullary myeloid sarcoma
ICD-O code 9930/3

Table 1.5 Sites of occurrence for head and neck extramedullary plasmacytomas

<table>
<thead>
<tr>
<th>Site</th>
<th>Frequency</th>
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<tbody>
<tr>
<td>Nasal cavity</td>
<td>28%</td>
</tr>
<tr>
<td>Paranasal sinuses</td>
<td>22%</td>
</tr>
<tr>
<td>Nasopharynx</td>
<td>22%</td>
</tr>
<tr>
<td>Tonsil</td>
<td>7%</td>
</tr>
<tr>
<td>Larynx</td>
<td>5%</td>
</tr>
<tr>
<td>Pharynx</td>
<td>5%</td>
</tr>
<tr>
<td>Soft palate</td>
<td>3%</td>
</tr>
<tr>
<td>Salivary gland</td>
<td>2%</td>
</tr>
<tr>
<td>Thyroid</td>
<td>1%</td>
</tr>
<tr>
<td>Tongue</td>
<td>1%</td>
</tr>
<tr>
<td>Gingiva</td>
<td>1%</td>
</tr>
<tr>
<td>Cervical lymph node</td>
<td>1%</td>
</tr>
<tr>
<td>Miscellaneous sites, e.g.,</td>
<td>2%</td>
</tr>
<tr>
<td>trachea, subcutaneous tissue</td>
<td></td>
</tr>
</tbody>
</table>

Immunohistochemistry
Immunohistochemically, the plasma cells express cytoplasmic immunoglobulin with light chain restriction. CD20 is negative in most cases, and some cases express CD79a. PAX-5 is negative, while Oct-2 and Bob.1 are frequently positive. There is usually expression of CD38, CD138 and VS38, markers characteristically positive in but not specific for plasma cells. Epithelial membrane antigen is commonly positive, and rare cases can show cytokeratin immunoreactivity (often with a dot pattern). Leukocyte common antigen, CD31 or CD56 is sometimes positive.

Extramedullary plasmacytoma. A Poorly-differentiated plasmacytoma. Note mononuclear and multinucleated neoplastic plasma cells with prominent nucleoli. B Immunoperoxidase staining of plasmacytoma shows monoclonal kappa staining in cytoplasm of neoplastic plasma cells. C Immunoperoxidase staining shows absence of staining for lambda light chains in neoplastic plasma cells.
Extramedullary myeloid sarcoma, also known as granulocytic sarcoma, is a tumour mass of myeloblasts or immature myeloid cells occurring outside the bone marrow or bone. It can precede, co-exist with or follow the presentation of acute myeloid leukaemia. It can also arise as a clastic transformation of an underlying chronic myeloproliferative disease or myelodysplastic syndrome.

The most common sites for occurrence of extramedullary myeloid sarcoma are lymph node and skin, but involvement of the nasal cavity and paranasal sinuses has also been reported [1701,2204]. The tumour mass comprises diffuse sheets of blast cells, which often show a single file pattern of infiltration in some areas. The blast cells have round or ovoid nuclei, very fine chromatin, small but distinct nucleoli, and a small to moderate amount of lightly eosinophilic cytoplasm. There can be better-differentiated cells with eosinophilic cytoplasmic granules. Intermingled eosinophilic myelocytes and metamyelocytes, if present, can provide an additional clue to the diagnosis. Giemsa-stained touch preparations are excellent for identification of cytoplasmic azurophilic granules as well as Auer rods, if present. Not uncommonly, extramedullary myeloid sarcoma is misdiagnosed as malignant lymphoma.

**Immunohistochemistry**

The tumour cells show chloroacetate esterase activity in approximately 75% of cases. Immunohistochemically, they express various myeloid markers (such as myeloperoxidase, CD13, CD33, CD117, CD68/KP1, neutrophil elastase and lysozyme), with myeloperoxidase being most sensitive and specific. Myeloid sarcoma with monocytic differentiation shows a myeloperoxidase-, CD68/PGM1+ immunophenotype. The pan-T marker CD43 is commonly expressed and may lead to a misdiagnosis of T-cell lymphoma.

**Histiocytic sarcoma**

ICD-O code 9755/3

Histiocytic sarcoma, defined as a malignant proliferation of cells showing morphologic and immunophenotypic features of mature tissue histiocytes, is a rare tumour that can occasionally present in the nasal cavity [2043]. The large pleomorphic tumour cells have eccentrically-located round, ovoid, indented or grooved nuclei, and abundant eosinophilic cytoplasm that may show fine vacuolation. Phagocytosis is rare. Histologic distinction from large cell lymphoma is difficult, except that the cytoplasm tends to be voluminous and eosinophilic. The diagnosis depends on the demonstration of histiocytic differentiation (granular staining for CD68 and lysozyme), in the absence of expression of pan-B markers (e.g. CD19, CD20, CD22, CD79a), pan-T markers (e.g. CD3), myeloid markers (e.g. MPO), Langerhans cell marker CD1a, and follicular dendritic cell markers (e.g. CD21, CD35). Since CD68 or lysozyme per se is not totally specific for histiocytic lineage, it is preferable to demonstrate additional haematolymphoid markers such as LCA/CD45, CD4, CD43 or CD163 to confirm the diagnosis. The frequent expression of CD43 may lead to a misdiagnosis of T-cell lymphoma. A small proportion of cases can express S100 protein.

**Langerhans cell histiocytosis**

ICD-O code 9751/1

Langerhans cell histiocytosis may occasionally present with nasal obstruction due to facial bone involvement [1183]. For details see Chapters 4 on tumours of the oral cavity and oropharynx and 7 on tumours of the ear.

**Juvenile xanthogranuloma**

This histiocytic proliferation may mimic tumours of Langerhans cells. It commonly presents as skin nodules in infants and children, but rare extracutaneous cases involving the nasal cavity and paranasal sinuses have also been reported [568, 2245]. Some of the histiocytes have foamy cytoplasm, and frequently there are scattered Touton giant cells and spindly cells within the infiltrate of nonde- script mononuclear cells. The histiocytes in juvenile xanthogranuloma express CD68 and factor XIIa. In contrast to Langerhans cells, they are negative for S100 protein and CD1a.

**Rosai-Dorfman disease**

Rosai-Dorfman disease (sinus histiocytosis with massive lymphadenopathy) is an uncommon reactive condition of unknown etiology, characterized by proliferation of distinctive histiocytes that usually exhibit emperipolesis of lymphocytes. The tumour masses can mimic lymphoma or other malignancies both clinically and histologically. The patients are commonly children or young adults who present with massively enlarged...
cervical lymph nodes. Extranasal involvement is frequent (about 40% of cases), especially the upper aerodigestive tract [791,923,1378,2760]. The patients with upper aerodigestive tract disease present with nasal obstruction, sinusitis, epistaxis, facial pain or saddle nose deformity.

Histologically, low magnification examination reveals alternating pale and dark-staining areas. The pale areas show proliferation of a distinctive type of very large histiocytes with round nuclei, distinct nucleoli, abundant light-staining cytoplasm, and indistinct cell borders. There are typically many admixed plasma cells. The characteristic histiocytes are usually much larger than the typical histiocytes with round nuclei, distinct nucleoli, abundant light-staining cytoplasm, and indistinct cell borders. The lesion may be mistaken for rhinoscleroma. In some cases, a proportion of histiocytes can exhibit atypical or irregular nuclei, and may lead to a misdiagnosis of malignancy such as histiocytic sarcoma or melanoma. The dark areas consist of large aggregates of plasma cells and small lymphocytes. Fibrosis is usually a prominent feature in extranasal disease, and the fibrotic bands can impart a nodular appearance. Together with spindling of some of the cells, the histologic features may strongly mimic those of fibrohistiocytic tumours or inflammatory pseudotumours. Immunohistochemically, the large histiocytes co-express S100 protein and the histiocytic marker CD68, and are negative for CD1a.

Most cases are treated by excision alone, with steroid, radiotherapy and chemotherapy having been given in a minority of cases. In general, the prognosis is excellent, most patients being free of disease or with stable disease. However, some patients may develop recurrent disease in the original site or other body sites.
Neuroectodermal tumours

Ewing sarcoma (EWS) / Primitive neuroectodermal tumour (PNET)

Definition
A high-grade, primitive, round cell tumour of neuroectodermal phenotype. EWS and PNET represent a group of small round cell neoplasms with variable degrees of neuroectodermal differentiation, and are considered together in this section under the rubric of EWS/PNET.

ICD-O codes
Ewing sarcoma 9260/3
PNET 9364/3

Synonyms
Peripheral neuroepithelioma, peripheral neuroectodermal tumour, peripheral neuroblastoma

Epidemiology
Sinonasal Ewing sarcoma / primitive neuroectodermal tumour (EWS/PNET) is rare. This is mostly a tumour of children and young adults, with a peak in the 3rd decade [2211]. In children, approximately 20% of EWS/PNET occur in the head and neck, with about 20% of these arising in the sinonasal tract [2122]. On rare occasion, older adults may be affected [2611]. There is a very slight male predominance [2122].

Localization
Sinonasal EWS/PNET most commonly occurs in the maxillary sinus [1518] and nasal fossa [1424,2069].

Clinical features
Symptoms include pain, mass, and obstruction [2069]. The tumour can be polypoid when arising from the nasal cavity [2069]. Bony erosion may or may not be present [2069].

Macroscopy
EWS/PNET is a grey-white and glistening tumour with haemorrhage and often ulceration [2069]. It tends to be much smaller than that arising in other sites.

Tumour spread and staging
Intranasal tumours usually spread into...
Tumours of the nasal cavity and paranasal sinuses

Histopathology
The tumour is composed of densely distributed, uniform, small to medium sized, round cells with a high nuclear to cytoplasmic ratio and fine chromatin. Mitotic activity is high, and coagulative necrosis is common. Homer Wright rosettes are rare.

Immunoprofile
The immunophenotype includes reactivity for CD99 (MIC2, O13, HBA-71, p30/32, and 12E7), vimentin, and on occasion focally for keratins. Some cases express neural markers, such as synaptophysin, S100 protein, NSE, neurofilament protein, GFAP, and chromogranin. Fli-1 (one portion of the gene fusion product of EWS/FLI1) can be detected by immunohistochemistry.

Electron microscopy
Electron microscopy reveals, to a variable extent, interdigitating neuritic processes, neurofilaments, microtubules, neurosecretory granules and glycogen.

Differential diagnosis
The differential diagnoses include malignant melanoma, melanotic neuroectodermal tumour, rhabdomyosarcoma, sinonasal undifferentiated carcinoma, lymphoma, olfactory neuroblastoma, and pituitary adenoma.

Histogenesis
A pluripotential fetal neuroectodermal cell is considered the progenitor.

Somatic genetics
Most EWS/PNET have a characteristic t(11;22) with EWS/FLI1 juxtaposition or other translocations involving EWS.

Genetic susceptibility
Sinonasal EWS/PNET has been reported in association with retinoblastoma.

Prognosis and predictive factors
EWS/PNET has much better prognosis in the head and neck than in other anatomic sites. Size and stage are the most important prognostic factors. Tumours demonstrating the EWS/FLI1 fusion are reported to have a better prognosis than those with less common gene fusion types. With improvements in imaging techniques and multimodality treatment, a 5-year survival of 60-70% can be achieved.

Olfactory neuroblastoma
Definition
A malignant neuroectodermal tumour thought to originate from the olfactory membrane of the sinonasal tract.

Synonyms
Esthesioneuroblastoma, esthesioneurocytoma, esthesioneuroepithelioma, olfactory placode tumour.

ICD-O code
9522/3

Epidemiology
Olfactory neuroblastoma is an uncommon neoplasm representing approximately 2-3% of sinonasal tract tumours. The incidence has been estimated at 0.4 per million. Patients range in age from as young as 2 years to 90 years, and a bimodal age distribution has been noted in the 2nd and 6th decades of life. Both genders are affected equally. No racial predilection has been noted.

Etiology
There are no known etiologic agent(s) for human olfactory neuroblastoma. Injection of diethylnitrosamine in Syrian hamsters and N-nitrosopiperidine in rats has produced tumours histologically identical to human olfactory neuroblastoma.

Localization
The most common site of origin is in the upper nasal cavity in the region of the cribriform plate. Included in the areas of the proposed origin are Jacobson’s organ (vomeronasal organ), sphenopalatine (pterygoid palatine) ganglion, olfactory placode, and the ganglion of Loci (nervus terminalis). “Ectopic” origin in lower nasal cavity or within one of the paranasal sinuses may occur. Olfactory neuroblastoma may occur.

Fig. 1.86 Olfactory neuroblastoma. A Tumour lobules separated by a highly vascularized stroma. B Olfactory neuroblastoma accompanied by the hyperplasia of the olfactory epithelium.
Neuroectodermal tumours

on occasion present as an intracranial (frontal lobe) mass with involvement of the superior aspect of the cribriform plate or rarely, occur intracranially with no intranasal component [987].

Clinical features

**Signs and symptoms**
The main presenting symptoms are unilateral nasal obstruction (70%) and episistaxis (46%); less common manifestations include anosmia, headache, pain, excessive lacrimation and ocular disturbances. Typically, these tumours are slow-growing resulting in long-standing symptomatology, the mean delay between the appearance of the first symptom and the diagnosis being 6 months [626].

**Imaging**
The radiologic features include the presence of a "dumbbell-shaped" mass extending across the cribriform plate. The extent of disease is best determined by pre- and postcontrast MR imaging in which there is intense signal in T2-weighted images with marked enhancement of T1-weighted images after gadolinium injection [2815]. Details of bone erosion (lamina papyracea, cribriform plate and fovea ethmoidalis) are better demonstrated by CT scan. Calcifications producing a speckled pattern on radiographic studies can be identified. Angiographic studies disclose a hypervascular neoplasm.

**Macroscopy**
The gross appearance includes a glistening, mucosa-covered, soft, polypoid, often highly vascularized mass varying from a small nodule measuring less than 1 cm to a large mass filling the nasal cavity and extending into paranasal sinuses, orbit and/or cranial cavity.

**Histopathology**
Characteristically, the tumours are localized to the submucosa, growing in circumscribed lobules or nests separated by a richly vascularized fibrous stroma. Less often the tumour shows a diffuse growth pattern. The overwhelming majority of tumours are not associated with an in-situ component. The neoplastic cells have uniform, small round nuclei with scant cytoplasm, dispersed ("salt and pepper") coarse to fine nuclear chromatin and inconspicuous nucleoli. Nuclear pleomorphism, mitotic activity and necrosis are usually absent. However, in higher-grade tumours, nuclear pleomorphism with prominent nucleoli, increased mitotic activity and necrosis may be present. The cells do not have distinct borders and are surrounded by a neurofibrillary matrix, which corresponds to tangles of neuronal cell processes. Rosettes of the Homer

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Fig. 1.87 Olfactory neuroblastoma. A Lobules of tumour separated by fibrovascular septa. B The lobules of tumour are separated by dense fibrovascular tissue. A large pseudorosette (Homer Wright) shows a central area of neurofibrillary matrix. C A high grade olfactory neuroblastoma showing a true Flexner-Wintersteiner rosette and increased mitotic figures. D The "small blue round cell" neoplasm has scant cytoplasm surrounding variably hyperchromatic nuclei. Granular nuclear chromatin can be seen. Mitotic figures are noted in higher grade lesions.
Wright type (pseudorosettes) and Flexner-Wintersteiner type (true neural rosettes) can be identified in up to 30% and less than 5% of tumours, respectively. The Homer Wright pseudorosettes represent the presence of cells in an annular arrangement surrounding central neurofibrillary matrix; distinct cell membranes are not present. Flexner-Wintersteiner rosettes are gland-like structures in which the annular arrangement of cells includes the presence of a distinct cell membrane. Perivascular pseudorosettes can be seen but are of no diagnostic utility. Uncommon findings include stromal calcifications, ganglion cells, melanin-containing cells and divergent differentiation. The latter may include the presence of glandular (adenocarcinoma-like), squamous, teratomatous and rhabdomyoblastic differentiation (1096,1734,2404).

**Grading**
The microscopic grading (1159) includes four grades: Grade I is the most differentiated and includes lobular architecture with intercommunication of the neoplasm between lobules. The neoplastic cells are well-differentiated with uniform, small round nuclei with scant cytoplasm, dispersed (“salt and pepper”) nuclear chromatin and inconspicuous nucleoli. The cells do not have distinct borders; rather, the nuclei are surrounded by a neurofibrillary material suggesting cytoplasmic extension. Homer Wright rosettes are frequently seen. Varying amounts of calcification may be noted. Interlobular fibrous stroma is often extremely vascular. Mitotic activity and necrosis are absent.

Grade II tumours share many of the histologic features described for Grade I lesions but the neurofibrillary element is less well defined, and the neoplastic nuclei show increased pleomorphism. Scattered mitoses can be seen. Grade III tumours may retain a lobular architecture with a vascular stroma. These hypercellular tumours are characterized by cells that are more anaplastic, hyperchromatic, and have increased mitotic activity as compared to Grade I or II tumours. Necrosis is seen. The neurofibrillary component may be focally present, but is much less conspicuous as compared to Grades I or II tumours. Flexner-Wintersteiner rosettes are uncommon. Calcification is absent. Grade IV tumours may also retain the overall lobular architecture, but the neoplastic element is the most undifferentiated and anaplastic of all the histologic grades. The cellular infiltrate is characterized by pleomorphic nuclei often with prominent eosinophilic nucleoli and an indistinct cytoplasm. Necrosis is commonly seen and there is increased mitotic activity, including atypical mitoses. Rosettes are uncommon. The neurofibrillary component is generally absent. Calcification is absent. Of note is that in any given tumour there may be histologic diversity with mixed (overlapping) features. In general, the lower grade olfactory neuroblastomas are readily recognizable and diagnostic by light microscopy. Adjunct studies, particularly in the higher histologic grade tumours, may assist in the diagnosis. The advent of immunohistochemistry has diminished the role of histochemical stains, but silver stains such as Bodian, Grimelius and Churukian-Schenk may still be of assistance.

**Immunoprofile**
The most consistently expressed marker is neuron specific enolase (NSE). Reactivity is also present in a majority of cases for synaptophysin, neurofilament protein (NFP), class III beta-tubulin, and microtubule-associated protein. S-100

<table>
<thead>
<tr>
<th>Microscopic Features</th>
<th>Grade 1</th>
<th>Grade 2</th>
<th>Grade 3</th>
<th>Grade 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Architecture</td>
<td>Lobular</td>
<td>Lobular</td>
<td>Lobular</td>
<td>Lobular</td>
</tr>
<tr>
<td>Pleomorphism</td>
<td>Absent to Slight</td>
<td>Present</td>
<td>Prominent</td>
<td>Marked</td>
</tr>
<tr>
<td>Neurofibrillary matrix</td>
<td>Prominent</td>
<td>Present</td>
<td>May be present</td>
<td>Absent</td>
</tr>
<tr>
<td>Rosettes</td>
<td>Present*</td>
<td>Present*</td>
<td>May be present**</td>
<td>May be present**</td>
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<tr>
<td>Mitoses</td>
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<td>Present</td>
<td>Prominent</td>
<td>Marked</td>
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<td>Necrosis</td>
<td>Absent</td>
<td>Absent</td>
<td>Present</td>
<td>Prominent</td>
</tr>
<tr>
<td>Glands</td>
<td>May be present</td>
<td>May be present</td>
<td>May be present</td>
<td>May be present</td>
</tr>
<tr>
<td>Calcification</td>
<td>Variable</td>
<td>Variable</td>
<td>Absent</td>
<td>Absent</td>
</tr>
</tbody>
</table>

NF-neurofibrillary; *Homer Wright rosettes (pseudorosettes); **Flexner-Wintersteiner rosettes (true neural rosettes)

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Fig. 1.88 Olfactory neuroblastoma. A A true Flexner-Wintersteiner rosette is surrounded by intermediate sized cells with scant cytoplasm and prominent nucleoli. A mitotic figure is present. B Immunostaining shows strong staining for synaptophysin. C Immunostaining shows the characteristic S100 protein+ sustentacular cells wrapping around the tumour islands.
protein staining typically is limited to the sustentacular cells situated along the periphery of the neoplastic lobules, although such cells may be sparse in the higher-grade tumours. In addition, immunoreactivity may be present for chromogranin, glial fibrillary acidic protein (GFAP), and Leu-7.

Cytokeratin is usually negative, but some cases can show some positive cells. Epithelial markers, including epithelial membrane antigen (EMA) and carcinoembryonic antigen (CEA) are absent. Leucocyte common antigen (LCA), HMB-45, desmin and CD99 are absent. Proliferation marker studies using Ki-67 and MIB-1 have shown a high proliferative index of 10-50% and flow cytometric analysis shows frequent polyploidy/aneuploidy {2560,2682}. Studies on cytogenetic aberrations in olfactory neuroblastoma are sparse {2612,2521}. Studies have found partial gains of chromosome material on 8q, while the other findings are conflicting. Inclusion of olfactory neuroblastoma within the Ewing sarcoma family of tumours {2428} or the primitive neuroectodermal tumours (PNET) {1865} has been proposed {2467} because of the identification, in certain cases, of translocation t(11:22), which is regarded as specific molecular abnormality for Ewing sarcoma {575}. Recent studies using immunohistochemistry, fluorescent in situ hybridization and reverse transcriptase PCR have failed to confirm this translocation in olfactory neuroblastoma {72, 1384, 1709, 2001}. Therefore, olfactory neuroblastoma should be considered an entity distinct from PNET and the Ewing sarcoma family of tumours.

Differential diagnosis

The differential diagnosis of olfactory neuroblastoma includes the group of small round cell malignant neoplasms that can occur in the sinonasal tract, i.e., sinonasal undifferentiated carcinoma, lymphoma, rhabdomyosarcoma, mucosal malignant melanoma and neuroendocrine carcinomas. This discussion will be limited to neuroendocrine carcinomas; for the others the reader may refer to the specific sections detailing these specific tumour types. Neuroendocrine carcinomas (NEC) include, among different tumour types, the carcinoid tumour, atypical carcinoid tumour and small cell carcinoma. NEC of the sinonasal tract are extraordinarily rare, and in contrast to the larynx, the most common subtype is small cell carcinoma. By light microscopy, small cell carcinoma typically is a submucosal hypercellular proliferation growing in sheets, cords and ribbons; the distinct lobular pattern of olfactory neuroblastoma is absent. The cells are small and hyperchromatic with oval to spindle-shaped nuclei, absent nucleoli and minimal cytoplasm. Cellular pleomorphism, high nuclear to cytoplasmic ratio, high mitotic activity, confluent necrotic areas and individual cell necrosis are readily apparent as well as lymphovascular and perineural invasion. Characteristically, crush artifacts of the neoplastic cells are seen. Squamous cell foci may occasionally be present; glandular or ductal differentiation is rarely seen. Although uncommon, neural-type rosettes similar to those seen in olfactory neuroblastoma can be seen in association with small cell carcinoma. The overall light microscopic findings should allow for differentiating small cell carcinoma from olfactory neuroblastoma in most cases, but immunohistochemical evaluation may be required in some cases. The immunohistochemical profile of small cell carcinoma includes variable reactivity for cytokeratin, chromogranin, synaptophysin, neuron specific enolase (NSE), S-100 protein and thyroid transcription factor-1 (TTF-1). Cytokeratin reactivity may include a punctate paranuclear or globoid pattern. The tumour usually is negative for cytokeratin, and the positive cases do not show a punctate paranuclear or globoid pattern. In contrast to olfactory neuroblastoma, NSE reactivity in small cell carcinoma is more likely to be focal than diffusely positive, and the S100 protein staining, if present, is dispersed throughout the cellular proliferation and not limited to sustentacular cells. Olfactory neuroblastoma is also negative for TTF-1.

Genetics

Studies on cytogenetic aberrations in olfactory neuroblastoma are sparse {2612,2521}. Studies have found partial gains of chromosome material on 8q, while the other findings are conflicting. Inclusion of olfactory neuroblastoma within the Ewing sarcoma family of tumours {2428} or the primitive neuroectodermal tumours (PNET) {1865} has been proposed {2467} because of the identification, in certain cases, of translocation t(11:22), which is regarded as specific molecular abnormality for Ewing sarcoma {575}. Recent studies using immunohistochemistry, fluorescent in situ hybridization and reverse transcriptase PCR have failed to confirm this translocation in olfactory neuroblastoma {72, 1384, 1709,2001}. Therefore, olfactory neuroblastoma should be considered an entity distinct from PNET and the Ewing sarcoma family of tumours.
Histogenesis

Proposed sources of origin of olfactory neuroblastoma include Jacobson’s vomero-nasal organ, the sphenopalatine ganglion, the ectodermal olfactory placode, Loci’s ganglion, autonomic ganglia in the nasal mucosa, and the olfactory epithelium. While a neuronal – neural crest origin is supported by the presence of neurofilaments in olfactory neuroblastoma (2634), until recently (335), few arguments linked olfactory neuroblastoma directly to the olfactory epithelium. The olfactory neuroepithelium is a unique neurosensory organ because olfactory neurons are continuously replaced throughout adult life by new ones (941,942). Three types of cells are classically recognized in the olfactory epithelium: the basal cells, located against the basement membrane, the olfactory neurosensory cells, and the sustentacular supporting cells, the processes of which extend on the luminal surface. The globose basal cells constitute a stem cell compartment, which confers to this tissue its peculiar ability to regenerate not only physiologically but also when injured by trauma or environmental insults (1631,2690). The globose basal cells express (1747) neural cell adhesion molecule (NCAM) (513) and mammalian homologue of Drosophila achaete-scute (MASH) gene (958). These progenitor cells differentiate into olfactory neurosensory cells, which exhibit a progressive maturation from the basal membrane to the epithelial surface (1631,1884). Each layer can be characterized by specific olfactory- and neuron specific markers. Immature olfactory cells express (1631,2690) GAP43, a 24 kD membrane-associated protein kinase C involved in polyphosphoinositide turnover (197). As these cells mature, they send axons to the olfactory bulb and migrate towards the surface, they express olfactory marker protein (OMP) (1630) and NCAM, but not GAP43 (1631,1884,2690).

Recently, olfactory neuroblastomas were found to express HASH, the human homologue of the MASH gene (335), while staining negative for OMP. So far, HASH has only been demonstrated in medullary thyroid carcinoma and certain small cell lung carcinoma (111). Further indirect evidence that olfactory neuroblastoma originates from olfactory stem cells can be derived from transgenic mice in which, the SV40T oncogene was inserted under the OMP gene promoter region (2307): these mice did not develop olfactory neuroblastoma but adrenal and sympathetic ganglia neuroblastoma. Therefore, the currently available evidence links olfactory neuroblastoma with the basal progenitor cells of the olfactory epithelium.

Prognosis and predictive factors

Complete surgical eradication (craniofacial resection that includes removal of the cribriform plate) followed by full course radiotherapy is the treatment of choice [625,626,1777]. Limited success using chemotherapeutic modalities have been achieved for advanced unresectable tumours and/or for disseminated disease {2705}. High-dose chemotherapy, including platinum-based protocols and autologous bone marrow transplantation have resulted in long-term survival [634,1919,2064]. The overall 5-, 10- and 15-year survival rates have been reported to be 78%, 71% and 68%, respectively [634]. Initial multimodality therapy is associated with 5-year survival of 80% for low-grade tumours and 40% for high-grade tumours [1777]. The majority of the recurrences occur within the first two years [625]. The most frequent recurrence is local, with rates around 30%. Prognosis has traditionally been correlated to clinical staging with 5-year survival of 75-91%, 68-71% and 41-47% for Stage A, B and C tumours, respectively [663,1243]. More recently, complete tumour resection was found to be of more prognostic importance than clinical staging [1740].

Other factors purportedly implicated in prognosis include histologic grading, proliferation rate and ploidy. Histologically lower grade tumours (Grades I and II) have been reported to have a better 5-year survival than higher grade tumours (Grades III, IV) [1159]. High proliferation indices and high rate of ploidy/aneuploidy have been correlated with increased morbidity (i.e., tumour recurrence, metastasis) and mortality (i.e., decreased survival) [2560,2682]. The majority of tumours behave as locally aggressive lesions mainly involving adjacent structures (orbit and cranial cavity). Local recurrence and distant metastasis may occur years following the initial diagnosis. Approximately 15-70% of patients will experience local recurrence, 10-25% will have cervical lymph node metastasis, and approximately 10-60% will experience distant metastasis (131,663). The more common sites of metastases include lymph nodes, lungs, and bone. All histologic grades have the capacity to metastasize.

<table>
<thead>
<tr>
<th>Stage</th>
<th>Extent of Tumour</th>
<th>5-Year survival</th>
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</thead>
<tbody>
<tr>
<td>A</td>
<td>Tumour confined to the nasal cavity</td>
<td>75-91%</td>
</tr>
<tr>
<td>B</td>
<td>Tumour involves the nasal cavity plus one or more paranasal sinuses</td>
<td>68-71%</td>
</tr>
<tr>
<td>C</td>
<td>Extension of tumour beyond the sinonasal cavities</td>
<td>41-47%</td>
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Melanotic neuroectodermal tumour of infancy

**Definition**

Melanotic neuroectodermal tumour of infancy (MNTI) is a rare neoplasm of infants with a biphasic population of neuroblastic cells and pigmented epithelial cells.

**ICD-O code**

9363/0 [if benign]

**Synonyms**

Melanotic progonoma, retinal anlage tumour, melanotic ameloblastoma

**Epidemiology**

The tumour is very rare. It characteristically occurs in infants, with 80% of cases <6 months of age and 95% <1 year of age, with a 2:1 female predominance [1269].

**Localization**

More than 85% of patients have a mass involving craniofacial sites: maxilla (70%), mandible (10%), skull (10%), neurocranial dura or brain (1%). Occasionally other sites, such as the epi-
Neuroectodermal tumours
didymis (4%), skin (3%), uterus (1%), and mediastinum (1%) may be involved [1269,2026].

Clinical features
Patients present with a rapidly growing pigmented mass, which is usually located in the anterior alveolar ridge of the maxilla. The duration of symptoms ranges from 2 weeks to 5 months (mean 2 months) [1269]. Infrequently, there are elevated levels of vanilmandelic acid, which normalize following adequate therapy.

Macroscopy
The tumours range from 1-10 cm (mean, 3.5 cm), and are smooth, firm to hard, and grey to blue-black [1269].

Histopathology
This is a nonencapsulated mass composed of a dual population of small neuroblastic cells and larger melanin-containing epithelial cells in a vascularized dense fibrous stroma [1269,2026]. The epithelial cells show alveolar or tubular arrangement, and often surround nests of smaller neuroblastic cells. The latter possess small, round hyperchromatic nuclei and scant or fibrillary cytoplasm. The epithelial cells have larger, vesicular nuclei and abundant cytoplasm, most containing melanin granules. Mitoses and necrosis are rare or absent.

Immunoprofile
MNTI shows polyphenotypic expression of neural, melanocytic and epithelial markers, but without photoreceptor differentiation. Occasionally glial and rhabdomyoblastic differentiation may be seen. The larger cell (epithelial) component is immunoreactive for cytokeratin, HMB-45, vimentin, and sometimes epithelial membrane antigen [1269,2026]. Neuron-specific enolase, CD57/Leu-7 and dopamine-beta-hydroxylase are often positive in both the small neuroblastic cells and large cells. The small cells may express synaptophysin, glial fibrillary acidic protein focally, and desmin focally. The tumour cells are negative for chromogranin, desmin, CEA, retinol-binding protein, neurofilaments, alpha-fetoprotein, and S100 protein.

Electron microscopy
The small cells demonstrate neurosecretory granules and neuritic processes, and the large cells contain melanosomes and premelanosomes [517,571,1269].

Differential diagnosis
The differential diagnoses include alveolar rhabdomyosarcoma, malignant lymphoma, EWS/PNET, metastatic neuroblastoma, immature teratoma and malignant melanoma. Primary melanoma, especially mucosal, is extremely rare in infants, should show S100 protein immunoreactivity, and lacks epithelial markers. Neuroblastomas may rarely be pigmented, but lack the dual cell population and usually show diffuse immunoreactivity for neuroendocrine markers.

Histogenesis
A neural crest origin is proposed [145, 517,571,1232,1269,2026].

Prognosis and predictive factors
The treatment of choice is complete local excision [1269,1657,2092]. Radiotherapy and chemotherapy are to be avoided,
Tumours of the nasal cavity and paranasal sinuses

unless there is evidence of metastasis. Despite its rapid growth and tendency to destroy bone, MNTI pursues a benign clinical course in most cases [1269,2026]. However, if not totally excised, local recurrences occur frequently. About 7% of cases develop metastases to sites such as the lymph nodes, liver, bone, adrenal glands or soft tissue [2026]. The potential for recurrence or metastasis, however, cannot be predicted from the clinical or pathologic features.

**Mucosal malignant melanoma**

**Definition**
A malignant neoplasm derived from the melanocytes in the mucosa.

**ICD-O code** 8720/3

**Synonyms**
Melanosarcoma; melanoma

**Epidemiology**
Sinonasal mucosal malignant melanomas are rare, accounting for less than 1% of all melanomas [112,165], and <5% of all sinonasal tract neoplasms [205,2603]. Both genders are equally affected, without a race predilection, although an increased incidence has been suggested in Japanese patients. Malignant melanomas typically affect older individuals in the 5-8th decade with a peak incidence in the 7th decade [165,260,273,386,484,500,560,807,930,1076,2603].

**Etiology**
Formaldehyde exposure and tobacco smoking have been suggested as possible etiologic factors [260,273,1318,2603].

**Localization**
The nasal cavity is affected most frequently, followed by a combination of the nasal cavity and paranasal sinuses. Large tumours may involve multiple paranasal sinuses and present as extensive skull base tumours [260,273,386,484,500,560,807,930,1324,2603].

**Tumour spread and staging**
At presentation, 70-80% of cases are localized, 10-20% have regional lymph node and <10% have distant metastasis [112,273,386,1324,2603]. However, dur-
Neuroectodermal tumours

The course of disease, an additional 20% may develop nodal metastasis and 40-50% may develop distant metastasis to lungs, brain, bone and/or liver (273,386,484,500,1324,2603). There is currently no universally accepted staging system. However, the most common and prognostically significant staging system in use is: stage I- localized tumours, stage II- tumours with lymph node metastases, and stage III- tumours with distant metastasis (1076). Tumour thickness or depth of invasion cannot be accurately assessed due to the lack of a well-defined reference point for the surface in the respiratory mucosa, frequent ulceration, tissue fragmentation and poorly oriented specimen (273,386,2603).

Histopathology
The tumours are comprised of epithelioid, spindled, plasmacytoid, rhabdoid, and/or multinucleated tumour cells. The cells are generally medium to large-sized (260,273,386,484,1472,2603). They have a high nuclear to cytoplasmic ratio with pleomorphic nuclei containing prominent eosinophilic nucleoli and intranuclear cytoplasmic inclusions. Nuclear molding can be present. The cytoplasm is usually densely eosinophilic, and variably contains melanin pigment. Mitoses, including atypical forms, are frequent and easily identifiable. Vascular invasion and neurotropism may be identified in up to 40% of cases. An inflammatory infiltrate admixed with pigment-laden histiocytes is commonly identified within or adjacent to the tumour. Tumour cell necrosis is common, particularly in tumours displaying a peritheliomatous or pseudopapillary growth pattern. Other growth patterns include solid, alveolar or sarcomatoid. Intraepithelial melanocytic atypia (melanoma in-situ) is sometimes seen in the overlying epithelium (260,273,386,1472,1624,2603). The tumours usually invade the subepithelial tissue and frequently extend into the bone, cartilage or skeletal muscle.

Immunoprofile
Malignant melanoma expresses S100 protein and vimentin (1472,1661), and variably HMB45, tyrosinase, melan-A and microphthalmia transcription factor. Neuron specific enolase, CD117, CD99, synaptophysin, CD56, and CD57 have been reported to be occasionally positive, but epithelial membrane antigen, cytokeratins, and muscle markers are not expressed (260,273,484,1472,1661,2603).

Differential diagnosis
Sinonasal mucosal malignant melanoma may morphologically masquerade as a variety of benign and malignant neoplasms, such as “small blue round cell” neoplasms, pleomorphic neoplasms.
Tumours of the nasal cavity and paranasal sinuses (sinonasal undifferentiated carcinoma, anaplastic large cell lymphoma, angiosarcoma), or various sarcomas. Metastatic melanoma to the sinonasal tract, although highly uncommon, must always be excluded, as the prognosis is even poorer. Presence of intraepithelial atypical melanocytes favours primary melanoma.

**Histogenesis**
Melanocytes, distributed throughout the upper respiratory tract are considered the progenitor of primary sinonasal mucosal malignant melanoma.

**Genetic susceptibility**
Patients with sinonasal mucosal malignant melanoma do not seem to be part of dysplastic nevus syndrome or xeroderma pigmentosum kindreds.

**Prognosis and predictive factors**
Surgery is the cornerstone of therapy, although wide free margins of resection are difficult to achieve. Radiation therapy has a palliative role. Local recurrence is frequent (67%-92%), may be repeated, and is a harbinger of adverse prognosis. Most tumours progress to regional and distant metastasis resulting in poor 5-year disease-specific survival that may range from 17-47%. Other poor prognostic factors include advanced age, obstructive symptoms, tumour size >3 cm, location in paranasal sinuses and nasopharynx, vascular invasion into skeletal muscle and bone, high mitotic count, marked cellular pleomorphism and distant metastasis.

**Epidemiology**
Most patients present at birth, and 90% of cases are diagnosed by age of 2 years. There is no gender predilection.

**Localization**
The lesion is situated externally on or near the bridge of the nose in 60% of cases, within the nasal cavity in 30% of cases, and in both sites in 10% of cases. In the latter cases, communication of the intra- and extranasal components is through a defect in the nasal bone.

**Clinical features**
Extranasal heterotopic CNS tissue presents as a smooth noncompressible subcutaneous mass over the dorsum of the nose. The intranasal lesions usually present with nasal obstruction or nasal deformity. Heterotopic CNS tissue may occur at other sites, such as the paranasal sinuses, nasopharynx, pharynx, tongue, palate, tonsil and orbit, and may be referred to as “facial glioma”. One-third of pharyngeal heterotopic CNS tissues are associated with cleft palate or choanal stenosis.

---

**Fig. 1.94 Heterotopic central nervous system tissue.**

A. Mucosal glands are subtended by reactive glial tissue composed of neuropil separated by dense, more brightly eosinophilic fibrous connective tissue. Astrocytes are not seen. B. S-100 positivity in nuclei and cytoplasm of subepithelial glial cells. C. The left side demonstrates a number of "gemistocytic-type" astrocytes within glial tissue, the right image shows classic neuroglial tissue without significant fibrosis or inflammatory cells. D. Trichrome stain highlights the neural tissue red, while the reactive background fibrosis is blue (left). GFAP immunoreactivity is present in glial tissue, but not in the surrounding soft tissues (right).
A helpful clinical sign is the absence of expansion or pulsation of the mass following compression of the ipsilateral jugular vein (negative Furstenberg test), due to lack of connection of the mass with the CSF pathway. Importantly, radiographic imaging scans (CT and MRI) reveal a soft tissue mass without an intracranial component or bony defect in the floor of the anterior cranial fossa.

**Macrosopy**
The lesion appears as a polypoid, smooth, soft, grey tan, non-translucent mass with encephaloid features. It usually measures 1-3 cm.

**Histopathology**
The lesion is non-encapsulated, composed of large or small islands of glial tissue with evenly spaced astrocytes and interlacing bands of vascularized fibrous connective tissue. The glial tissue merges with the collagen of the stroma or dermis. Mitoses are absent. At times, the astrocyte nuclei may appear enlarged or multinucleated. Long-standing or recurrent lesions tend to contain a considerable amount of fibrous tissue. Neurons are rare or absent. Rarely, choroid plexus, ependyma-lined clefts and pigmented retinal epithelium are seen, especially those of the palate and nasopharynx.

The glial tissue can be confirmed by immunoreactivity for glial fibrillary acidic protein (GFAP) or S100 protein [607,1273,1323,1991,2851].

**Differential diagnosis**
The histologic differential diagnoses mainly include nasal encephalocele and, less frequently, a fibrosed nasal polyp. In contrast to heterotopic CNS tissue, encephaloceles are herniations of meningeal lined brain tissue that communicate with the intracranial ventricular system and subarachnoid space through a bony defect in the skull [1134]. Nasal encephalocele is composed of CNS tissue with easily found neurons. However, in nasal encephalocele of long-standing and in recurrences, the excessive fibrous tissue relative to the amount of glial cells and absence of neurons may make it impossible to distinguish from heterotopic CNS tissue.

Long-standing heterotopic CNS tissue may be mistaken for a fibrosed nasal polyp [1258]. The absence of glial tissue differentiates the latter from the former.

**Histogenesis**
It is a congenital malformation in which there is anterior displacement of mature cerebral tissue that has lost connection with the intracranial contents.

**Prognosis and predictive factors**
Adequate excision offers a cure in most cases, but incomplete excision can be accompanied by recurrence (15-30%). There is no local aggressive behaviour or malignant potential.

**Ectopic pituitary adenoma**
This lesion is described in Chapter 2 on tumours of the nasopharynx.
**Germ cell tumours**

*Malignant germ cell tumours and teratocarcinosarcoma* exhibiting histologic features similar to germ cell tumours of the gonads arise on rare occasions in the sinonasal tract. Immature teratomas and teratomas with malignant transformation are tumours of infancy and early childhood, whereas sinonasal yolk sac tumour and sinonasal teratocarcinosarcoma have only been documented in adults.

**Immature teratoma**

ICD-O code 9080/3

Immature teratomas are rare in the sinonasal tract and nasopharynx, and are composed of variable quantities of immature tissue elements, mostly neuroepithelial, that are interspersed with mature and immature tissues derived from the three embryonic germ layers. They are tumours of infancy and childhood (2317).

Immature teratomas tend to be either solid-nodular or solid-cystic, while mature teratomas are usually cystic. The tumour may contain cystic spaces lined by ciliated pseudostratified epithelium as well as primitive neuroepithelium with rosettes. Mitotic figures are frequently present in the immature areas; however, cellular atypia is not found. In infants and children, a teratoma with malignant transformation has to be excluded. In adults, thorough sampling of the specimen is mandatory to rule out teratocarcinosarcoma. Immature teratomas rarely behave in a malignant fashion (570).

**Sinonasal yolk sac tumour (endodermal sinus tumour)**

ICD-O code 9071/3

This is a tumour that has the histological features of embryonic yolk sac, indistinguishable from yolk sac tumour (endodermal sinus tumour) of the gonads. Only two cases have been reported to arise in the sinonasal tract (1623). Both patients were adults (aged 34 and 43 years). In one case, there was an admixed component of sinonasal nonkeratinizing carcinoma. The behaviour has been aggressive.

**Sinonasal teratocarcinosarcoma**

**Definition**

A complex malignant sinonasal neoplasm combining features of teratoma and carcinosarcoma. Benign and malignant epithelial, mesenchymal, and neural elements are typically present, including immature tissue with blastomatous features, while embryonal carcinoma, choriocarcinoma or seminoma is absent.

**Synonyms**

Malignant teratoma, blastoma, teratocarcinoma, teratoid carcinosarcoma

**Epidemiology**

Sinonasal teratocarcinosarcoma is very rare (755). Approximately 60 cases have been published (755,1042,1619,1970, 2339,2578,2749). Patients are exclusively adults, with age ranging from 18-79 years (mean 60 years). There is a marked male predominance (1042).

**Localization**

It almost exclusively arises in the ethmoid sinus and maxillary antrum. One tumour has been reported to arise in the roof of the nasopharynx and another from the dorsum of the tongue (1042).

**Clinical features**

Patients present with a short history of nasal obstruction and epistaxis (1042). Imaging studies reveal a nasal mass, occasionally accompanied by opacification of the paranasal sinuses. Bone destruction may be seen.

**Macroscopy**

Tumours are usually bulky, soft to rubbery, and red-tan to purple.

**Teratoma with malignant transformation**

ICD-O code 9084/3

Teratoma with malignant transformation is a neoplasm containing benign tissue elements of all three germinal layers and, in addition, a somatic malignancy. There is only a single reported case involving the sinonasal tract of a 13-month-old boy. The malignant component was a squamous cell carcinoma (1379). The tumour was locally aggressive and recurred after surgery. There was no further recurrence 2 years after chemotherapy (1379).

**Fig. 1.95** Yolk sac tumour of nasal cavity. Typical features of yolk sac tumour, with many hyaline globules.
Histopathology
There are multiple tissue types derived from two or three germ layers, exhibiting variable degrees of maturity. In addition, there are intermingled carcinomatous and sarcomatous components [755, 2319]. The epithelial component is usually made up of keratinizing and nonkeratinizing squamous epithelium, pseudodifferentiated columnar ciliated epithelium, and glandular structures lined by either cuboidal or columnar cells that may show mucous differentiation. Nests of immature squamous cells containing clear cells (fetal-looking) are a common finding and an important diagnostic clue [1042]. The carcinomatous component is usually glandular, but sometimes squamous. Neuroepithelial elements with rosettes and neuroblastoma-like areas are present in most instances. The mesenchymal areas range from immature tissues (such as cartilage) to sarcomas (such as rhabdomyosarcoma and fibrosarcoma). There may be a proliferation of small round cells that are difficult to classify.

Immunoprofile
The undifferentiated/primitive component often shows positive immunoreactivity for CD99 and occasionally synaptophysin and S-100 protein [1970]. The spindle cell component is consistently positive for vimentin, and sometimes desmin, myoglobin, and glial fibrillary acidic protein. The neuroepithelial component is positive for neuron-specific enolase and occasionally chromogranin, alfa-fetoprotein, and cytokeratin. The epithelial component is positive for cytokeratins, epithelial membrane antigen, and occasionally S-100 protein and glial fibrillary acidic protein.

Differential diagnosis
Inadequate sampling may lead to erroneous diagnoses of olfactory neuroblastoma, squamous cell carcinoma, undifferentiated carcinoma, adenocarcinoma, malignant salivary gland-type tumours and adenosquamous carcinoma [1042].

Histogenesis
The tumour is unlikely to be of germ cell origin, but probably arises from a primitive cell in the olfactory/sinonasal membrane that not only reproduces the neuroectodermal features of olfactory neuroblastoma, but also has the capacity to differentiate into divergent types of somatic cells [1970].

Prognosis and predictive factors
Teratocarcinosarcomas are highly malignant. They are locally aggressive, rapidly invading soft tissue and bone as well as the orbit and cranial cavities. They also have the potential to metastasize to regional lymph nodes and distant sites, mainly the lungs. The average survival is less than 2 years, with 60% of the patients not surviving beyond 3 years [1042]. Recurrences usually appear within 3 years.

Mature teratoma
Teratoma is the principal benign germ cell tumour of the sinonasal region and shows histologic features similar to its
counterparts in the gonads and in other extragonadal locations.

**Definition**
Tumour composed of a variety of mature tissues that are foreign to the site of occurrence. There are typically tissues derived from two or three germ layers.

**ICD-O code** 9080/0

**Synonyms**
Teratoid tumour, benign teratoma.

**Epidemiology**
Teratomas of the head and neck account for only 6% of all teratomas (2558). Mature teratomas in the sinonasal tract are even more uncommon (955). Most cases occur in neonates and older infants, with equal sex distribution (955,1737). Stillbirth, prematurity, fetal malpresentation, dystocia, and maternal polyhydramnios are frequent associations.

**Localization**
In the sinonasal tract, the maxillary antrum and nasal cavity are affected more often than the sphenoid sinus (1036,1408,1778,1805,2312). The nasopharynx can also be the primary site of involvement.

**Clinical features**
Facial deformity, nasal obstruction, and a nasal mass are common manifestations. Occasional calcifications may be seen on imaging (955,1805). Teratomas may be associated with other skull deformities, anencephaly, hemicrania, and palatal fissures [8].

**Macroscopy**
The tumours are usually cystic, but can be solid or multilocular. They are commonly encapsulated masses that measure up to 7 cm.

**Histopathology**
Teratomas are composed of variable admixtures of mature skin, skin appendages, fat, glial tissue, smooth muscle, cartilage, bone, minor salivary glands, respiratory epithelium and gastrointestinal epithelium. Neural tissues are seen more often in sinonasal teratomas than in teratomas of other sites. Although the variegated histologic appearance of mature teratomas is usu-
Histologically, dermoid cysts are lined with mature keratinizing squamous epithelium and frequently contain cutaneous appendages in the cyst wall. This lesion is differentiated from a teratoma by the limited variety of tissue types and the absence of endodermal components. Epidermal inclusion cysts may resemble dermoid cysts but do not contain adnexa and occur predominantly in adults [99,582,2622].

**Histogenesis**
The most likely explanation for the ontogeny of dermoid cysts is the retention of ectodermal tissue along the lines of closure at junctions of bones, soft tissues, and embryonic membranes.

**Prognosis and predictive factors**
Dermoid cysts are treated by complete surgical excision. Recurrence is uncommon (<7%) [582,2891].
Secondary tumours

Definition
Tumours that involve the nasal cavity and paranasal sinuses that originate from, but are not in continuity with, primary malignant neoplasms of other sites. Leukaemias and lymphomas are excluded.

Epidemiology
Metastases to the nasal cavity and paranasal sinuses are rare (1300,2085) and may occur in any age group. In a review of 82 cases, the median age of patients with metastatic tumours at diagnosis was 57 years (range 3 months to 76 years), and about 60% were males (202).

Localization
The distribution of tumour among the paranasal sinuses and the most frequent tumour types to metastasize to these sites are shown in the Table 1.8 and 1.9. In 10-15% of cases, the metastases are limited to the nasal cavity.

Clinical features
Metastases to the sinonasal tract are haematogenous. They may be solitary or multifocal and ordinarily produce symptoms indistinguishable from those of a primary tumour. These include nasal obstruction, headache, facial pain, visual disturbances, exophthalmos, facial swelling, cranial nerve deficits and epistaxis (especially metastatic renal and thyroid carcinomas). In some instances, the metastasis may be the first manifestation of an otherwise clinically occult carcinoma, usually renal cell carcinoma.

Prognosis and predictive factors
Although the eventual outcome is usually poor, prognosis depends, in part, on whether the sinonasal metastasis is isolated or part of widespread disseminated disease. If the metastasis to the nasal cavity and sinuses is localized and treated aggressively, the average survival following discovery of the metastasis may be as long as 20-30 months (1300).

<table>
<thead>
<tr>
<th>Table 1.8</th>
<th>Most frequent sites of primary tumours that metastasize to the paranasal sinuses*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Primary Tumour</td>
<td>Frequency</td>
</tr>
<tr>
<td>Kidney</td>
<td>40%</td>
</tr>
<tr>
<td>Lung</td>
<td>9%</td>
</tr>
<tr>
<td>Breast</td>
<td>8%</td>
</tr>
<tr>
<td>Thyroid</td>
<td>8%</td>
</tr>
<tr>
<td>Prostate</td>
<td>7%</td>
</tr>
<tr>
<td>Miscellaneous</td>
<td>28%</td>
</tr>
</tbody>
</table>

*Data derived from reference (2085)

<table>
<thead>
<tr>
<th>Table 1.9</th>
<th>Distribution of 168 tumours metastatic to paranasal sinuses*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sinus</td>
<td>Frequency</td>
</tr>
<tr>
<td>Maxillary</td>
<td>33%</td>
</tr>
<tr>
<td>Sphenoid</td>
<td>22%</td>
</tr>
<tr>
<td>Ethmoid</td>
<td>14%</td>
</tr>
<tr>
<td>Frontal</td>
<td>9%</td>
</tr>
<tr>
<td>Multiple sinuses</td>
<td>22%</td>
</tr>
</tbody>
</table>

*Data based on reference (2085)

Fig. 1.99 Metastatic carcinoma. A Renal cell carcinoma metastatic to the maxillary sinus showing clear cells (due to accumulation of glycogen in the cytoplasm) and prominent sinusoidal vascularity. B Secondary prostatic adenocarcinoma. A malignant epithelial proliferation is identified within the large, patulous vessels in the sinonasal tract mucosa. C (Same case as B). The metastatic nature of the tumour (left) was confirmed when the prostate specific antigen (PSA) was strongly and diffusely immunoreactive in the cytoplasm of tumour cells within vascular spaces (right).
CHAPTER 2

Tumours of the Nasopharynx

A wide variety of tumours can arise in the nasopharynx, but it is nasopharyngeal carcinoma that has fascinated generations of oncologists, pathologists, scientists and epidemiologists. It shows marked geographic differences, with highest incidence rates in Southern Chinese. In some endemic areas, the incidence has declined by about 30% over the past two decades, suggesting that environmental or lifestyle factors may play a major role and that the disease is, to some extent, preventable. Nasopharyngeal carcinoma shows a very strong association with Epstein-Barr virus (EBV) infection, irrespective of the ethnic origin of the patients. This association has pioneered a new paradigm of utilizing viral serological tests for the diagnosis of cancer and for screening in high-risk populations. Nasopharyngeal carcinoma is generally responsive to radiation therapy, and the clinical outcome has greatly improved over the years, due to refinements in staging and to improved therapy protocols.

The unusual and often deceptive histological features of nasopharyngeal carcinoma have generated controversies over the nature of the tumour and still pose a challenge to surgical pathologists. There have possibly been more names invented for the various histological subtypes of nasopharyngeal carcinoma than any other tumour type. The WHO classification presented in this book is expected to become the world-wide standard reference.
WHO histological classification of tumours of the nasopharynx

Malignant epithelial tumours

Nasopharyngeal carcinoma

- Nonkeratinizing carcinoma 8072/3
- Keratinizing squamous cell carcinoma 8071/3
- Basaloid squamous cell carcinoma 8083/3
- Nasopharyngeal papillary adenocarcinoma 8260/3
- Salivary gland-type carcinomas

Soft tissue neoplasms

- Nasopharyngeal angiofibroma 9160/0

Haematolymphoid tumours

- Hodgkin lymphoma
- Diffuse large B-cell lymphoma 9680/3
- Extramedullary plasmacytoma 9734/3
- Follicular dendritic cell sarcoma/tumour 9758/3

Nasopharyngeal papillary adenocarcinoma 8260/3

Benign epithelial tumours

- Hairy polyp
- Schneiderian-type papilloma 8121/0
- Squamous papilloma 8050/0
- Ectopic pituitary adenoma 8272/0
- Salivary gland anlage tumour
- Craniopharyngioma 9350/1

Tumours of bone and cartilage

- Chordoma 9370/3

Salivary gland anlage tumour

Craniopharyngioma

Distant metastasis

- M- MX Distant metastasis cannot be assessed
- M0 No distant metastasis
- M1 Distant metastasis

Stage Grouping

- Stage 0 Tis N0 M0
- Stage I T1 N0 M0
- Stage IIA T2a N0 M0
- Stage IIB T1 N1 M0
- Stage IIb T2a, T2b N1 M0
- Stage III T1 N2 M0
- Stage IV A T4 N0, N1, N2 M0
- Stage IVB Any T N3 M0
- Stage IVC Any T Any N M1

TNM classification of carcinomas of the nasopharynx

TNM classification

<table>
<thead>
<tr>
<th>T-Primary Tumour</th>
<th>N-regional lymph nodes**</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX Primary tumour cannot be assessed</td>
<td>NX Regional lymph nodes cannot be assessed</td>
</tr>
<tr>
<td>T0 No evidence of primary tumour</td>
<td>N0 No regional lymph node metastasis</td>
</tr>
<tr>
<td>T1 Tumour confined to nasopharynx</td>
<td>N1 Unilateral*** metastasis in lymph node(s), &lt;6 cm in greatest dimension, above the supraclavicular fossa</td>
</tr>
<tr>
<td>T2 Tumour extends to soft tissues</td>
<td>N2 Bilateral metastasis in lymph node(s), &lt;6 cm in greatest dimension, above the supraclavicular fossa</td>
</tr>
<tr>
<td>T2a Tumour extends to oropharynx and/or nasal cavity without parapharyngeal extension*</td>
<td>N3 Metastasis in lymph node(s), &gt;6 cm and/or in the supraclavicular fossa</td>
</tr>
<tr>
<td>T2b Tumour with parapharyngeal extension*</td>
<td>N3a &gt;6 cm in dimension</td>
</tr>
<tr>
<td>T3 Tumour invades bony structures and/or paranasal sinuses</td>
<td>N3b in the supraclavicular fossa#</td>
</tr>
<tr>
<td>T4 Tumour with intracraniad extension and/or involvement of cranial nerves, infratemporal fossa, hypopharynx, orbit, or masticator space</td>
<td></td>
</tr>
</tbody>
</table>

Distant metastasis

- M- MX Distant metastasis cannot be assessed
- M0 No distant metastasis
- M1 Distant metastasis

Stage Grouping

- Stage 0 Tis N0 M0
- Stage I T1 N0 M0
- Stage IIA T2a N0 M0
- Stage IIB T1 N1 M0
- Stage IIb T2a N1 M0
- Stage III T1 N2 M0
- Stage IV A T4 N0, N1, N2 M0
- Stage IVB Any T N3 M0
- Stage IVC Any T Any N M1

*Parapharyngeal extension denotes postero-lateral infiltration of tumour beyond the pharyngobasilar fascia.
** The regional lymph nodes are the cervical nodes.
*** Midline nodes are considered ipsilateral nodes.
# Supraclavicular fossa is the triangular region defined by 3 points: the superior margin of the sternal end of the clavicle, the superior margin of the lateral end of the clavicle, the point where the neck meets the shoulder. This includes caudal portions of Levels IV and V.

1 (847,2418).
2 A help desk for specific questions about the TNM classification is available at www.uicc.org/index.php?id=508.
The most common type of nasopharyngeal tumour is nasopharyngeal carcinoma, which is remarkable for the striking geographic differences in its incidence as well as the near consistent association with the Epstein-Barr virus (EBV). Nasopharyngeal carcinoma is also the prototype of a family of morphologically distinctive tumours – the lymphoepithelial carcinomas – that can arise in a variety of sites, such as other head and neck mucosal sites, salivary gland, lung and thymus, albeit uncommonly. Interestingly, in contrast to nasopharyngeal carcinoma, lymphoepithelial carcinomas occurring in these sites usually show a strong association with EBV only in Asians, but not in Caucasians. Besides nasopharyngeal carcinoma, a broad range of neoplasms can arise in the nasopharynx, from epithelial to lymphoid, mesenchymal and neurogenic. Rarely, tumours derived from embryonic remnants either entrapped in their normal pathway of ascent or descent (ectopic pituitary tumour, cranioopharyngioma) or dissociated from their normal regulatory influences (germ cell tumour) can occur. Since the nasopharynx is in close proximity to many different anatomical structures, tumours arising in the latter sites can also present clinically as a nasopharyngeal mass, for example, chordoma arising in the clivus.

Anatomy
The nasopharynx is the narrow tubular passage behind the nasal cavity. Its sloping roof and posterior wall are formed by the basi-sphenoid, basisphenoid, and the first cervical vertebra. Anteriorly, it communicates with the nasal cavity via the choanae. The orifices of Eustachian tubes are in the lateral walls, and each is shielded superiorly and posteriorly by a comma-shaped elevation called the torus tubarius. Immediately above and behind the torus tubarius is a pharyngeal recess called the fossa of Rosenmüller. The nasopharynx tapers inferiorly, and continues as the oropharynx from the level of the soft palate.

The nasopharynx constitutes part of the Waldeyer ring. Histologically, its mucosa is covered by respiratory-type ciliated epithelium, but variable amounts of squamous epithelium are common. The mucosa exhibits invaginations, forming crypts that abut the underlying stroma. The stroma is rich in lymphoid tissue that often includes reactive lymphoid follicles. The surface or crypt epithelium is commonly infiltrated by many small lymphoid cells, which expand and disrupt the epithelium to produce a reticulated pattern. Some seromucinous glands are present, but they are not as abundant as in the nasal mucosa.

Clinical features
Diagnostic procedures
Various imaging techniques, such as computed tomography and magnetic resonance imaging, are helpful for detection of the presence of a tumour, as well as in precise delineation of the extent of disease. Endoscopic examination with directed biopsy is the key in obtaining materials for a definitive histological diagnosis.

Tumour staging
The TNM staging system for nasopharyngeal tumours (see preceding section) is only applicable for epithelial tumours, and in fact has been developed specifically for nasopharyngeal carcinoma. For lymphomas, the Ann Arbor staging system is recommended (947).

Classification of nasopharyngeal carcinomas
In the 1978 WHO classification, three histological subtypes of nasopharyngeal
carcinoma were recognized: squamous cell carcinoma (WHO type 1), nonkeratinizing carcinoma (WHO type 2), and undifferentiated carcinoma (WHO type 3) (2320). In the 1991 WHO classification, the squamous cell carcinoma subtype (keratinizing squamous cell carcinoma) was retained, while the last two subtypes in the previous classification were combined under a single category of “nonkeratinizing carcinoma”, which was further subdivided as being “differentiated” or “undifferentiated”; lymphoepithelioma-like carcinoma was considered a morphologic variant of undifferentiated carcinoma (2317). The use of numerical designation of WHO types 1, 2 and 3 was eliminated. The wide ranging reported figures on the frequencies of various subtypes indicate that the boundaries between the categories are not always clear (such as less well differentiated forms of keratinizing squamous cell carcinoma versus nonkeratinizing carcinoma, and nonkeratinizing carcinoma versus undifferentiated carcinoma), sampling error is a significant problem due to the small size of the biopsies, and intra-and inter-observer reproducibility of the classification is sub-optimal (323,2318, 2497,2735). In fact, squamous cell carcinoma and nonkeratinizing carcinoma have been viewed by some investigators as being merely variants of a fairly homogeneous group of tumours (2318,2577). Notwithstanding these problems, the proportion of keratinizing squamous cell carcinoma among all nasopharyngeal carcinomas is probably higher in low-incidence compared with high-incidence areas.

The current WHO classification maintains the terminology of the 1991 classification, with the addition of one category: basaloid squamous cell carcinoma.

Fig. 2.1A  Global incidence rates of cancer of the nasopharynx (all ages) in males. Age-standardized rates (ASR, world standard population) per 100,000 population and year. From: Globocan 2002 (http://www-depdb.iarc.fr/globocan/GLOBOframe.htm).
Nasopharyngeal carcinoma

Definition
A carcinoma arising in the nasopharyngeal mucosa that shows light microscopic or ultrastructural evidence of squamous differentiation. It encompasses squamous cell carcinoma, nonkeratinizing carcinoma (differentiated or undifferentiated) and basaloid squamous cell carcinoma. Adenocarcinoma and salivary gland-type carcinoma are excluded.

ICD-O codes
Nonkeratinizing carcinoma
8072/3
Keratinizing squamous cell carcinoma
8071/3
Basaloid squamous cell carcinoma
8083/3

Synonyms
Lymphoepithelioma, lymphoepithelioma-like carcinoma, lymphoepithelial carcinoma, Schmincke type lymphoepithelioma, Regaud type lymphoepithelioma, transitional cell carcinoma, intermediate cell carcinoma, anaplastic carcinoma, undifferentiated carcinoma with lymphoid stroma, vesicular nucleus cell carcinoma, squamous cell carcinoma (WHO-1), nonkeratinizing carcinoma (WHO-2), undifferentiated carcinoma (WHO-3).

Epidemiology
Global incidence and mortality
Nasopharyngeal carcinoma (NPC) shows a distinct racial and geographical distribution and a multifactorial etiology. Globally, there were approximately 65,000 new cases and 38,000 deaths in the year 2000 (730). While rare in most parts of the world (onset rates commonly <1 per 105, or 0.6% of all cancers), there are certain populations for which the incidence is considerably higher, notably native and foreign-born Chinese, Southeast Asians (e.g. in Thailand, Philippines, and Vietnam), North Africans (e.g. in Algeria and Morocco), as well as native peoples of the Arctic region (e.g. in Canada and Alaska). Within these populations, there is a remarkable heterogeneity among ethnic lines (2872). The highest incidence of NPC has long been observed in Hong Kong, where 1 in 40 men develop NPC before the age of 75 years (1981).

Age and sex distribution
In high-risk groups, NPC incidence rises after the age of 30 years and peaks at 40-60 years, and thereafter declines (730). The age distribution is similar in males and females, although rates in men are commonly 2-3-fold those observed in women (1981).

Migration
In general, populations that migrate from high to low risk areas retain much of the elevated risk [304] seen in their country of origin, although this, and the extent to which the risk diminishes in successive generations, varies according to ethnicity. Such heterogeneity may be associated with several factors, possibly acting in combination - the degree of genetic predisposition, and the prevalence of certain risk factors related to lifestyle upon migration.

Time trends
Recent trends in NPC incidence in high-risk countries reveal convincing evidence of a decline in rates since the mid-1970s in Hong Kong (1446). The speed of the decline points to the role of changing environmental risk factors. Rates in low-risk areas are, in view of the rarity of the disease, subject to a great deal of random variation, and trends are often difficult to interpret. In U.S. Whites and in England, rates are low and in slow decline. The evolution of trends in U.S. Blacks is unclear.

Etiology
The specific geographical and demographic distribution of nasopharyngeal carcinoma (NPC), the time trends, and patterns observed in migrants reflect the interplay of genetic susceptibility, infection by Epstein-Barr virus (EBV) and environmental factors (dietary and non-dietary) in disease causation.
**Epstein-Barr virus**

The near constant association of EBV with NPC, irrespective of ethnic background, indicates a probable oncogenic role of the virus in the genesis of this tumour \(1166,2107\). The evidence includes: (1) raised levels of antibodies, especially IgA, against EBV (most commonly viral capsid antigen and early antigen) in most patients with NPC compared with normal controls and patients with other cancer types; (2) higher titers of IgA antibodies against EBV in patients with large tumour bulk; (3) presence of EBV DNA or RNA in practically all tumour cells; (4) presence of EBV in a clonal epimodal form, indicating that the virus has entered the tumour cell before clonal expansion; (5) presence of EBV in the precursor lesion of NPC, but not in the normal nasopharyngeal epithelium. The evidence was considered sufficient to classify EBV as carcinogenic by the International Agency for Research on Cancer (IARC) in 1997 \(1166\). Nevertheless, it is likely that EBV infection takes place relatively late in the oncogenic process \(1893\).

The EBV infection in NPC exhibits the type II latency pattern, that is, expression of EBV nuclear antigen-1 (EBNA-1) and latent membrane protein-1 (LMP-1), but not the immunogenic EBNA2-6. EBV encoded early RNAs (EBERs) are expressed in abundance. BZLF (Zebra) protein, which is expressed in lytic infection by EBV, is not detected. LMP-1, a viral protein with transforming properties, can induce epidermal hyperplasia, inhibit squamous differentiation, upregulate the adhesion molecule ICAM-1 and CD40, activate nuclear factor-κB (NF-κB), and induce expression of epidermal growth factor receptor.

**Environmental factors**

**Diet**

In high incidence regions, high levels of volatile nitrosamines in preserved food have been implicated as the putative carcinogen for NPC development \(2063\). In the 1960s, it was proposed that the increased incidence of NPC among Hong Kong boat dwellers compared to house dwellers may have been due to their staple diet of salted fish \(1108\). From case-control studies on Chinese, consumption of Cantonese-style salted fish in the weaning period carries odds ratios ranging from 2.75 \(2869,2870,2913\). Animals fed salted fish may develop nasopharyngeal tumours \(1142,2871,2913\). Other preserved or fermented foods in high incidence regions, consumed during weaning and early childhood, have also been incriminated as risk factors \(2869,2903\). The importance of exposure in early life is supported by two studies showing that low-risk ethnic groups born in high-risk areas also have higher risk of NPC \(1212,1213\). In low incidence regions like Northern China, the consumption of salted fish still carries an adjusted relative risk as high as 5.6 \(304\).

**Other environmental risk factors**

Other purported risk factors include cigarette smoking \(1900,2868,2875\), occupational exposure to smoke, chemical fumes and dusts, formaldehyde exposure \(74,1090\), and prior radiation exposure \(403\).

**Localization**

The most common site of origin is the lateral wall of the nasopharynx, especially the fossa of Rosenmüller, followed by the superior posterior wall. Cervical lymph node metastasis is a common occurrence.

**Clinical features**

**Signs and symptoms**

About half of the patients have multiple symptoms, but 10% are asymptomatic. Painless enlargement of upper cervical lymph node(s) is the most common presenting feature. Nearly half of the patients complain of nasal symptom(s), particularly blood stained post-nasal drip. Symptoms related to Eustachian tube obstruction (such as serous otitis media) also commonly occur. Headache and symptoms related to cranial nerve...
involvement are features of more advanced disease. In endemic areas, NPC is an important underlying malignancy in patients presenting with dermatomyositis, being found in 12% of patients [2005] although only 1% of patients with NPC have dermatomyositis [2573].

**Imaging**

Magnetic resonance (MR) is the study of choice for assessing the loco-regional extent because of its superior sensitivity and multiplanar capability. Although CT is useful in depicting cortical bone erosion, MR is superior in revealing soft tissue infiltration and intracranial extension [442,1880] as well as marrow replacement permitting early recognition of bony involvement [442,1880]. Systemic imaging workup for patients with high metastatic risk include X-ray/CT of chest, ultrasonography/CT of liver, isotope bone scan and positron emission tomography coupled with CT (PET-CT).

**Table 2.03** Structures involved by local infiltration of nasopharyngeal carcinoma at presentation. Source: Magnetic resonance studies of 308 patients from Pamela Youde Nethersole Eastern Hospital, Hong Kong.

<table>
<thead>
<tr>
<th>Structures involved</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adjacent soft tissues</td>
<td></td>
</tr>
<tr>
<td>Nasal cavity</td>
<td>87%</td>
</tr>
<tr>
<td>Oropharyngeal wall, soft palate</td>
<td>21%</td>
</tr>
<tr>
<td>Parapharyngeal space, carotid space</td>
<td>68%</td>
</tr>
<tr>
<td>Pterygoid muscle (medial, lateral)</td>
<td>48%</td>
</tr>
<tr>
<td>Prevertebral muscle</td>
<td>19%</td>
</tr>
<tr>
<td>Bony erosion / paranasal sinus</td>
<td></td>
</tr>
<tr>
<td>Nasal septum</td>
<td>3%</td>
</tr>
<tr>
<td>Pterygoid plate(s), pterygo-maxillary fissure, pterygo-palatine fossa</td>
<td>27%</td>
</tr>
<tr>
<td>Maxillary antrum</td>
<td>4%</td>
</tr>
<tr>
<td>Ethmoid sinus</td>
<td>6%</td>
</tr>
<tr>
<td>Sphenoid sinus, sphenoid bone, foramina lacerum, ovale, rotundum</td>
<td>38%</td>
</tr>
<tr>
<td>Clivus</td>
<td>41%</td>
</tr>
<tr>
<td>Petrous bone, petro-occipital fissure</td>
<td>19%</td>
</tr>
<tr>
<td>Jugular foramen, hypoglossal canal</td>
<td>4%</td>
</tr>
<tr>
<td>Pituitary fossa / gland</td>
<td>3%</td>
</tr>
<tr>
<td>Extensive/ intracranial extension</td>
<td></td>
</tr>
<tr>
<td>Cavernous sinus</td>
<td>16%</td>
</tr>
<tr>
<td>Cerebrum, meninges, cisterns</td>
<td>4%</td>
</tr>
<tr>
<td>Infratemporal fossa</td>
<td>9%</td>
</tr>
<tr>
<td>Orbit, orbital fissure(s)</td>
<td>4%</td>
</tr>
<tr>
<td>Hypopharynx</td>
<td>2%</td>
</tr>
</tbody>
</table>

**Serological studies**

Positive serology against Epstein-Barr virus (EBV) is found in close to 100% of patients with non-keratinizing NPC (976). IgA against viral capsid antigen (VCA) and IgG/IgA against early antigens (EA) are the most extensively used diagnostic tool, with the reported detection rates for NPC varying from 69-93%. Newer antibody tests based on recombinant EBV antigens such as EBV nuclear antigens (EBNA), membrane antigen (MA), thymidine kinase (TK), DNA polymerase (DP), ribonucleotide reductase (RR), DNAase, and Z transactivator protein (Zta) have shown promise when used in combination [378,537].

Another approach is to test for elevated levels of circulating EBV DNA or RNA (BamH1-W, EBERs or EBNA1) by quantitative PCR (Q-PCR) in the plasma or serum, with reported sensitivity in NPC up to 96% [35,1514,1549,2346].

**Staging**

The current TNM staging system [947, 2418] is customized for NPC, as the natural

**Macroscopy**

The tumour can appear as a smooth bulge in the mucosa, a discrete raised nodule with or without surface ulceration, or a frankly infiltrative fungating mass. Sometimes no grossly visible lesion is seen.

**Tumour spread and staging**

**Tumour spread**

NPC is notorious for its highly malignant behaviour, with extensive loco-regional infiltration, early lymphatic spread, and disproportionately high incidence of haematogenous dissemination. Erosion of skull base and paranasal sinuses, intracranial spread (via eroded bone or basal foramina), infiltration of cranial nerves, and extension to more distant structures (infratemporal fossa, orbit, hypopharynx) occur as tumour invasion advances.

With the rich lymphatic plexus in the nasopharynx, lymphatic spread occurs early in the course of disease. In patients staged by MR, about 20% of patients have no enlarged nodes, and about half have retropharyngeal node involvement [2314]. The jugulo-digastric node is by far the most common palpable node at presentation, and involvement of the posterior cervical chain is more frequent than with other head and neck cancers. The strong association between the topographic level of lymphatic extension and the increased incidence of distant failure reflects that haematogenous dissemination occurs mainly via the draining of the lymphatic trunks at the lower end of the jugular chain into the great vessels. The most common sites of haematogenous deposits are, in descending order of frequency, bone, liver, lung, and distant nodes [2575].

**Relevant diagnostic procedures**

All patients should have complete physical examination and endoscopic examination of the nasopharyngeal region. Biopsies are taken from the gross lesions. In the absence of a gross lesion, multiple biopsies should be taken from the lateral, superior and posterior walls of the nasopharynx for patients with high suspicion of NPC.
behaviour and therapeutic considerations of NPC are so uniquely different from other head and neck cancers. With accumulation of supporting data from different countries \cite{448,490,1061,1119,1444,1592,1966}, there is little doubt that the current staging system is superior to the past systems, both in terms of improved predictive accuracy and more balanced stage distribution.

**Nonkeratinizing carcinoma**

**Histopathology**

The biopsies vary in appearance from the presence of a frank tumour with surface ulceration to subtle involvement of the mucosa beneath an intact surface epithelium \cite{2316,2318,2735}. The tumour comprises solid sheets, irregular islands, dyscohesive sheets and trabeculae of carcinoma intimately intermingled with variable numbers of lymphocytes and plasma cells. Subclassification into the undifferentiated and differentiated subtypes is optional, since their distinction is of no clinical or prognostic significance, and different areas of the same tumour or different biopsies taken at different time intervals from the same patient may exhibit features of one or the other subtype. When both subtypes are seen in a specimen, the tumour may be classified according to the prominent subtype, or as nonkeratinizing carcinoma with features of both subtypes.

The undifferentiated subtype, which is more common, is characterized by syncytial-appearing large tumour cells with indistinct cell borders, round to oval vesicular nuclei, and large central nucleoli. The cells often appear crowded or even overlapping. Sometimes, the nuclei

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**Fig. 2.6** Nasopharyngeal nonkeratinizing carcinoma. Tumour islands are obvious in the lymphoid stroma.

**Fig. 2.7** Nasopharyngeal nonkeratinizing carcinoma. This example is accompanied by an abundant desmoplastic stroma.

**Fig. 2.8** Nasopharyngeal nonkeratinizing carcinoma. A This example of differentiated subtype is characterized by sheets of tumour separated by a dense infiltrate of lymphocytes and plasma cells. B Tumour island in a lymphoid cell-rich stroma. Some lymphocytes are also seen within the tumour. C This tumour shows an uncommon trabecular growth pattern.

**Fig. 2.9** Cytological spectrum of nasopharyngeal nonkeratinizing carcinoma, undifferentiated subtype. A The cells exhibit a syncytial quality, and possess vesicular nuclei, prominent nucleoli and amphophilic cytoplasm. B The syncytial-appearing cells have vesicular nuclei, distinct nucleoli and lightly eosinophilic cytoplasm. There are some intermingled lymphocytes. C Focally, there can be cells with more distinct cell borders and a moderate amount of eosinophilic cytoplasm.
Nasopharyngeal carcinoma can be chromatin-rich rather than vesicular. The scant cytoplasm is either amphophilic or eosinophilic. There can be small foci of primitive squamous differentiation, where groups of tumour cells exhibit a slightly greater amount of lightly eosinophilic cytoplasm and slightly more distinct cell borders.

The differentiated subtype differs from the undifferentiated subtype in showing cellular stratification and pavementing, often with a plexiform growth, reminiscent of transitional cell carcinoma of the bladder [2317]. The tumour cells show fairly well-defined cell borders and sometimes vague intercellular bridges, and there may exceptionally be occasional keratinized cells. Compared with the undifferentiated subtype, the cells are often slightly smaller, the nuclear-cytoplasmic ratio is lower, the nuclei can be more chromatin-rich, and nucleoli are usually not as prominent.

A desmoplastic stroma is uncommon. Areas of coagulative necrosis are sometimes present, and can be extensive. The density of lymphocytes and plasma cells is highly variable. At one extreme, there are no or few lymphocytes within the tumour islands, although some lymphoid cells are present in between, which probably merely represent the native lymphoid tissue in the nasopharyngeal mucosa. At the other extreme, abundant lymphocytes and plasma cells infiltrate the tumour islands, breaking them up into tiny clusters or single cells and obscuring the epithelial nature of the tumour; the term “lymphoepithelial carcinoma” may be applied for such cases. In metastatic sites, the lymphocyte density in the tumour may or may not be maintained. In some cases, scattered epithelioid granulomas are present, and may be so prominent as to mask the small islands of carcinoma [404]. Many admixed eosinophils are seen in about one-fourth of cases [830,1463,1555]. Some cases show a prominent infiltrate of neutrophils even in the absence of ulceration.

There are a number of inconstant features. The carcinoma cells can assume a plump or slender spindle shape focally or extensively, with formation of streaming fascicles. The nucleoli of the spindly cells are often not as prominent as the syncytial-appearing ones. In some cases, isolated or groups of tumour cells may appear shrunken, with dark smudged nuclei and dense amphophilic or eosinophilic cytoplasm; it is unclear whether such changes reflect a degenerative phenomenon in a subpopulation of tumour cells, or a biopsy artefact. There can be Pagetoid spread of the carcinoma into the surface or crypt epithelium. In approximately one-tenth of cases, there are interspersed spherical amyloid globules [2084]. The amyloid globules are usually smaller than a tumour cell, and can be present intracellularly (sometimes causing indentation of the tumour cell nucleus), scattered among the carcinoma cells, or in the adjacent stroma. They are derived from keratins, and are probably of tumour origin. In the uncommon papillary variant, there are papillary fronds comprising delicate stromal cores covered by stratified tumour cells morphologically no different from those of the usual nonkeratinizing nasopharyngeal carcinoma. Rare cases can show cyto-

Table 2.04 Frequency of histological subtypes of nasopharyngeal carcinoma.

<table>
<thead>
<tr>
<th>Current WHO classification</th>
<th>High incidence population</th>
<th>Intermediate incidence</th>
<th>Low incidence population</th>
</tr>
</thead>
<tbody>
<tr>
<td>Keratinizing squamous cell carcinoma</td>
<td>1%</td>
<td>17%</td>
<td>8%</td>
</tr>
<tr>
<td>Nonkeratinizing carcinoma</td>
<td>99%</td>
<td>83%</td>
<td>92%</td>
</tr>
<tr>
<td>- Undifferentiated</td>
<td>92%</td>
<td>42%</td>
<td>76%</td>
</tr>
<tr>
<td>- Differentiated</td>
<td>7%</td>
<td>41%</td>
<td></td>
</tr>
<tr>
<td>Basaloid squamous cell carcinoma</td>
<td>&lt;0.2%</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

NA = Not available; *Queen Elizabeth Hospital, 2001-2003
plasmic clear cell change, but this is such an uncommon feature that the alternative diagnosis of lymphoma or salivary gland-type carcinoma should always be considered. Exceptionally, there is accumulation of extracellular edema fluid or mucosubstance, breaking up the tumour islands to produce a complex reticulated pattern. Nasopharyngeal carcinoma may contain intracytoplasmic mucin in very rare cells. It has also rarely been reported to occur in combination with a component of adenocarcinoma (1200, 1389). Nasopharyngeal carcinoma may present initially with cervical lymph node metastases. The lymph nodes can be involved extensively or subtly (such as submergence of the tumour in the lymphocyte-rich paracortex). The tumour takes the form of islands and strands, being intermingled with variable numbers of lymphocytes, plasma cells and eosinophils. Some tumour cells can resemble Reed-Sternberg cells or lacunar cells. Coupled with a dense lymphoid infiltrate, a misdiagnosis of Hodgkin or non-Hodgkin lymphoma is sometimes made (330,1470). A desmoplastic stroma may be present. In approximately one fifth of cases, there are epithelioid granulomas, and in half of these cases, the granulomas show caseous necrosis (1470). Nasopharyngeal carcinoma may also metastasize as a wholly or partly cystic lesion containing necrotic material.

**Immunoprofile**

Practically all tumour cells show strong staining for pan-cytokeratin (AE1/AE3, MNF-116); this uniform staining contrasts with the usually focal staining observed in undifferentiated carcinomas of other sites, such as the lung and thyroid. The staining for high molecular weight cytokeratins (such as cytokeratin 5/6, 34ßE12) is strong, and staining for low molecular weight cytokeratins (such as CAM5.2) is often weaker and sometimes patchy. Cytokeratins 7 and 20 are both negative (801). In undifferentiated nonkeratinizing carcinoma, the cytokeratin immunostain highlights the scanty wisps of cytoplasm that wrap around the large nucleus and extend outward as short narrow processes. As a result of the cell nests being broken up by infiltrating lymphocytes, a distinctive reticulated or meshwork pattern is produced. In differentiated nonkeratinizing carcinoma, the tumour cells, with a broader rim of cytoplasm, are obviously polygonal on immunostaining for cytokeratin. Immunoreactivity for epithelial membrane antigen in nasopharyngeal carcinoma is often only focal (816). In most cases, the tumour exhibits strong nuclear staining for p63, a basal cell marker that normally highlights the basal and parabasal cells of the overlying stratified squamous epithelium. The lymphoid cells represent a mixture of T cells and B cells, usually with the former predominating, especially within and around the tumour islands (854,883,1070,1962,2912). At least a proportion of the T cells are activated cytotoxic cells. The plasma cells are polyclonal. There are variable numbers of scattered S100

![Fig. 2.11 Nasopharyngeal nonkeratinizing carcinoma with papillary architecture.](image1)

A The tumour forms exophytic papillae with fibrovascular cores. B The lining cells show features of differentiated nonkeratinizing carcinoma.

![Fig. 2.12 Nasopharyngeal nonkeratinizing carcinoma, undifferentiated subtype.](image2)

A In-situ hybridization for EBER shows that all tumour cells exhibit nuclear labeling. B Immunostaining for pan-cytokeratin highlights the surface epithelium as well and irregular clusters and sheets of positive cells (carcinoma) in the stroma. C Immunostaining for cytokeratin usually reveals a meshwork pattern of staining.
protein-positive dendritic cells. Some studies have reported the following features to be associated with a better prognosis: high density of dendritic cells; high number of infiltrating lymphocytes; and low number of granzyme B-positive cytotoxic cells [854,883,1903,1962,2912].

**Epstein-Barr virus detection**

Nonkeratinizing nasopharyngeal carcinoma is associated with Epstein-Barr virus (EBV) in practically 100% of cases, irrespective of the ethnic background of the patient. EBV latent membrane protein-1 (LMP1) is usually positive in only 30-40% of cases, and the immunostaining is often patchy and weak, and thus is not a reliable method to demonstrate the presence of EBV [16,961,1988,2061]. The simplest and most reliable way to demonstrate EBV is in-situ hybridization for EBV encoded early RNA (EBER), which is present in abundance in cells latently infected by EBV. Practically all the tumour cells should show nuclear labelling [1137,1157,1176,2061,2233,2638,2684]. In-situ hybridization for EBER can aid in the diagnosis of nasopharyngeal carcinoma if there are difficulties in distinguishing between carcinoma and reactive epithelial atypia. A positive result also strongly suggests a nasopharyngeal origin (although not entirely specific) for a metastatic nonkeratinizing carcinoma of unknown primary. On the other hand, fine needle aspiration cytological examination of enlarged cervical lymph nodes is invaluable in reaching a diagnosis of metastatic nasopharyngeal carcinoma, either for initial diagnosis or staging [380,1355]. The aspirate smears show, in a background of lymphocytes and plasma cells, irregular clusters of large cells with overlapping vesicular nuclei and large nucleoli. The cytoplasm of these cells is often fragile and barely visible. There are commonly many naked nuclei [1760]. The presence of dispersed large tumour cells among the lymphoid cells may result in a pattern strongly reminiscent of Hodgkin lymphoma [1355]. The diagnosis can be readily confirmed by immunostaining for cytokeratin and in-situ hybridization for EBER either on the cell smears or cell block preparations.

**Electron microscopy.**

Although squamous differentiation is primitive or not evident in most cases of nasopharyngeal carcinoma at the light microscopic level, there is usually convincing evidence of squamous differentiation at the ultrastructural level. At least some carcinoma cells contain small bundles of tonofilaments or tonofibrils, in addition to well-formed desmosomes [1470, 1513,2082,2568].

**Differential diagnosis**

Crush artefacts are common in nasopharyngeal biopsies, making it difficult to determine whether the observed distorted cells represent carcinoma or merely lymphoid cells. Such biopsies should be scrutinized in the better-preserved areas for tumour cell clusters. If there are uncertainties, immunostaining for cytokeratin is of great help in reaching a diagnosis of nasopharyngeal carcinoma. In the non-neoplastic nasopharyngeal mucosa, cytokeratin immunostaining highlights the sharply delineated surface and crypt epithelium, with no positive cells in the stroma other than those in seromucinous glands. Mucosa involved by nasopharyngeal carcinoma typically shows irregular clusters of cytokeratin-positive cells in the stroma.

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**Fig. 2.13 Metastatic nasopharyngeal carcinoma in lymph node.** A Fine needle aspiration smear shows tight clusters of tumour cells among small lymphocytes. B In histological sections, examination under medium magnification often reveals areas where cohesive tumour growth is evident. C The epithelial nature of the tumour is readily confirmed by immunostaining for cytokeratin, whereby a meshwork pattern of staining is often observed.

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**Fig. 2.14 Nasopharyngeal mucosa.** A Germinal centre cells mimicking nasopharyngeal carcinoma. B Nasopharyngeal lymphoid hyperplasia mimicking nasopharyngeal carcinoma. In the left field, the venule with no obvious lumen can also be mistaken for a cluster of carcinoma cells.
A number of benign cellular changes can mimic nonkeratinizing carcinoma. (1) Clusters of germinal centre cells may be mistaken for carcinoma because of the presence of large cells with vesicular nuclei and the absence of a well-defined mantle zone. The identification of admixed centrocytes (smaller “atypical” cells with irregular-shaped or angulated nuclei) and tingible-body macrophages points toward the lymphoid nature of the large cells, which can be confirmed by immunostaining (leucocyte common antigen positive, cytokeratin negative) [2317]. (2) A tangentially sectioned crypt harbouring cells with reactive changes that include nuclear enlargement can produce a pattern simulating an island of carcinoma lying in a lymphoid cell-rich stroma. In contrast to nasopharyngeal carcinoma, the nuclei are not as large and thus not so crowded, and the nucleoli are not as prominent. A negative in-situ hybridization for EBER would render a diagnosis of nasopharyngeal carcinoma most unlikely [2318]. (3) The nasopharyngeal mucosa can sometimes exhibit reactive lymphoid hyperplasia, accompanied by an increased number of immunoblasts in the lymphoid stroma, raising a suspicion for carcinoma. In contrast to the latter, the large cells are non-cohesive and have well-defined amphophilic cytoplasm. The diagnosis can be confirmed by a lack of cytokeratin immunoreactivity as well as positive immunostaining for lymphoid markers in the large cells [323]. (4) The lymphoid tissue-associated venules lined by plump endothelial cells with vesicular nuclei may be mistaken for clusters of carcinoma cells. The presence of distinct basement membrane around the groups of cells, lack of large nucleoli, and negative staining for cytokeroloi, and negative staining for cytokeratin would be against the diagnosis of carcinoma. Distinction between nonkeratinizing carcinoma and large cell lymphoma can at times be difficult. In the nasopharyngeal mucosa or metastatic deposit in lymph node, dispersed growth of the carcinoma cells and accompanying eosinophil infiltration may lead to a misdiagnosis of Hodgkin lymphoma [330,394,2880]. Features favouring a diagnosis of carcinoma include the presence of cohesive cell groups in some foci (best appreciated at medium magnification) and the generally poorly defined cell borders; the diagnosis can be readily confirmed by immunostaining for cytokeratin. Nasopharyngeal carcinoma with marked cellular spindling can mimic a high-grade sarcoma. In most cases, the diagnosis can be reached by identifying in some foci a component of typical nasopharyngeal carcinoma, and can be further confirmed by cytokeratin immunoreactivity.

Post-treatment biopsies
After treatment by radiation therapy, it may take weeks (up to 10 weeks) for the nasopharyngeal carcinoma to disappear histologically [1401]. The radiated carcinoma cells usually show evidence of radiation injury in the form of enlarged and bizarre nuclei, accompanied by an increased amount of cytoplasm that is often finely vacuolated. If biopsy is positive, repeat biopsies should be taken every two weeks – remission is defined by two subsequent negative biopsies [1401,1402,1886]. Radiation-induced changes in the normal nasopharyngeal mucosa can be mistaken for malignancy. The surface or crypt epithelium can exhibit enlarged, hyperchromatic or even bizarre nuclei, but such changes can be recognized to be benign because they are limited to some but not all cells (random cytoplastic atypia) and the normal nuclear-cytoplasmic ratio is maintained. Mucosal epithelial atypia usually does not persist beyond one year. If there are uncertainties as to whether the atypical cells represent residual carcinoma or irradiated normal cells, positive in-situ hybridization for EBER would strongly favour the former interpretation. There can also be bizarre stromal cells (radiation fibroblasts) with large smudged nuclei or large vesicular nuclei with prominent nucleoli; these atypical cells can persist for many years. These cells can be distinguished from residual or recurrent carcinoma by their occurrence as single cells and by the amphophilia of the cytoplasm. The stroma frequently contains ectatic blood vessels showing variable degrees of radiation injury such as enlarged prominent endothelial cells and abundant fibrinoid deposits. Some patients with nasopharyngeal carcinoma develop local recurrence. The nasopharyngeal biopsies should be interpreted in the same way as for patients without a prior history of nasopharyngeal carcinoma. The recurrence can be morphologically identical to the original tumour, or may show a slightly greater degree of squamous dif-
differentiation. Some recurrences, especially those occurring after a long interval (>5 years), may represent new primaries rather than a genuine relapse of the original tumours [1445]. Radiation-induced tumours in the nasopharynx typically develop after a long latency period, and usually take the form of keratinizing squamous cell carcinomas or sarcomas (especially osteosarcomas) [403,600].

**Keratinizing squamous cell carcinoma**  
*Histopathology*

This is an invasive carcinoma showing obvious squamous differentiation at the light microscopic level, in the form of intercellular bridges and/or keratinization over most of the tumour, morphologically similar to keratinizing squamous cell carcinomas occurring in other head and neck mucosal sites [2317]. The degree of differentiation can be further graded as: well differentiated (most common), moderately differentiated and poorly differentiated. The tumour typically grows in the form of irregular islands, accompanied by an abundant desmoplastic stroma infiltrated by variable numbers of lymphocytes, plasma cells, neutrophils and eosinophils [1555]. The tumour cells are polygonal and stratified. The cell borders are distinct and separated by intercellular bridges. The cells in the centres of the islands or facing the surface often show a greater amount of eosinophilic glassy cytoplasm, sometimes with identifiable cytoplasmic tonofibrils, indicative of cellular keratinization. Occasionally keratin pearls are formed [2735]. The nuclei often show hyperchromasia, and the degree of nuclear pleomorphism ranges from mild to marked. The surface epithelium is frequently involved, apparently representing carcinoma in-situ.

Keratinizing squamous cell carcinoma can arise de novo or as a radiation-associated carcinoma occurring many years after radiation therapy for nonkeratinizing nasopharyngeal carcinoma [403,2316,2735]. Compared with nonkeratinizing carcinoma, keratinizing squamous cell carcinoma shows a greater propensity for locally advanced tumour growth (76% versus 55%) [2136] and a lower propensity for lymph node metastasis (29% versus 70%) [1859]. While some studies suggest that this subtype of nasopharyngeal carcinoma has lower responsiveness to radiation therapy and a worse prognosis compared with nonkeratinizing carcinoma [1122,1859,2136,2318], others have not found this subtype to differ in biological behaviour [363,778].

**Immunoprofile and Epstein-Barr virus detection**

Keratinizing squamous cell carcinoma shows immunoreactivity for pan-cytokeratin, high molecular-weight cytokeratin, and focally epithelial membrane antigen. For radiation-induced keratinizing squamous cell carcinoma, there is no association with EBV [403]. However, for de novo keratinizing squamous cell carcinomas, data on the EBV status are conflicting. In general, the patients have lower or negative IgA titres against EBV compared with nonkeratinizing carcinomas [1486,1860,2549]. Molecular studies of EBV in the tumour tissues have yielded conflicting results. Summarizing the literature, it appears that EBV is almost always positive in areas endemic for nasopharyngeal carcinoma, EBV is often positive in intermediate incidence areas, while EBV is positive in only a proportion of cases in low incidence areas [405,580,599,961,1124,1125,1157,1176,1262,1885,1892,1988,2894]. Keratinizing squamous cell carcinomas tend to carry lower copy numbers of EBV compared with nonkeratinizing carcinomas [2108]. On in situ hybridization, the nuclear signals of EBER are usually confined to the less differentiated cells (basal cells that surround the individual tumour islands), but not in the cells showing obvious squamous differentiation.

The role of human papillomavirus in keratinizing squamous cell carcinoma remains uncertain [1125].

**Differential diagnosis**

The frank invasive growth, nuclear atypia and obvious squamous differentiation usually permit a straight-forward diagnosis of keratinizing squamous cell carcinoma to be made. However, in some cases, particularly those arising after radiation therapy for nonkeratinizing nasopharyngeal carcinoma, distinction between a very well differentiated keratinizing squamous cell carcinoma and squamous metaplasia/hyperplasia can be extremely difficult, since the nuclear atypia can be very subtle and focal, and invasion may not be obvious in the former. Assessment of invasion is further hampered by the abundant fibrinous deposits in the stroma related to prior radiation, and the usual desmoplastic stroma may be lacking. To arrive at a definitive diagnosis, sometimes multiple biopsies are required to identify convincing stromal invasion as well as focal mild nuclear atypia.

**Basaloid squamous cell carcinoma**

Several cases of basaloid squamous cell carcinoma, morphologically identical to the same tumour more commonly occur-
ring in other head and neck sites (See chapter on hypopharynx, larynx and trachea for details), have been reported to occur as primary tumours of the nasopharynx \{116,117,1790,1997,2714\}. Among the 6 cases with information, the M:F ratio is 2:1, and patients’ ages ranged from 27-79 years (mean 55 years). Four cases had stage T3 or T4 disease; and two had lymph node metastasis. None had distant metastasis at presentation. On follow-up, three patients had no evidence of disease at 34-52 months; three were alive with disease at 19-46 months. The tumour appears to show a lower clinical aggressiveness compared with basaloid squamous cell carcinoma occurring in other head and neck sites. Among 4 cases tested for EBV, all three Asian cases were positive, while one Caucasian case were negative \{1790,2714\}.

**Precursor lesions**
In biopsies of nasopharyngeal carcinoma, an in-situ or intraepithelial component is identified in only 3-8% of cases, but it is often difficult to determine whether the invasive carcinoma has originated from the overlying in situ carcinoma or has merely invaded the surface epithelium \{364,1504,1989,2852,2911\}. Pure nasopharyngeal carcinoma in-situ, as confirmed by multiple biopsies to rule out an invasive component, is very rare \{419,1989,2911\}. These findings suggest that most nasopharyngeal carcinomas do not originate from nasopharyngeal carcinoma in-situ, or the evolution from the latter to the former occurs over a short time scale such that the latter is rarely detected.

Histologically, pure nasopharyngeal carcinoma-in-situ is characterized by atypical epithelial change confined to the surface or crypt epithelium, and lacking an invasive component. The epithelium is usually slightly thickened, and consists of cells with variable loss of polarity, nuclear enlargement, nuclear crowding and distinct nucleoli. Sometimes there can be scattered amyloid globules. Some attempts have been made to grade the spectrum of intraepithelial neoplastic changes (dysplasia/carcinoma-in-situ, or nasopharyngeal intraepithelial neoplasia) in the nasopharynx, but reproducibility and difficulties in recognizing the lower grade lesions remain an issue.

So far, all cases of nasopharyngeal carcinoma in-situ studied have been positive for EBV (EBER), confirming that EBV infection precedes the acquisition of invasiveness by nasopharyngeal carcinoma \{419,1971,2813\}. Analysis of the EBV termini shows the virus to be in a clonal form, providing indirect support for the clonality of the epithelial proliferation \{1989\}. Thus in situ hybridization for EBER may aid in the distinction between carcinoma-in-situ and non-specific reactive atypia of the nasopharyngeal epithelium.

There are only limited data on the natural history of untreated pure nasopharyngeal carcinoma in-situ (or dysplasia). A proportion of patients develop invasive cancer on follow-up \{1971,1989\}.

**Histogenesis**
Nasopharyngeal carcinoma arises from the surface or crypt epithelium of the nasopharyngeal mucosa. In some cases, the tumour appears to arise from the basal layers of the stratified squamous epithelium, a finding further supported by the strong immunoreactivity for p63 in both the tumour and normal basal cells.

**Somatic genetics**
Nasopharyngeal carcinoma (NPC) is believed to result from accumulation of multiple genetic alterations and Epstein-Barr virus (EBV) latent infection in the
nasopharyngeal epithelial cells [611, 1542]. EBV genome is detected in all undifferentiated NPC cells and in high-grade dysplastic lesions of the nasopharynx, but rarely found in the adjacent normal nasopharyngeal epithelial cells or in the low-grade dysplastic lesions [361, 1989]. The expression of EBV latent genes (e.g. EBNA1, LMP-1, LMP-2) may alter multiple signal transduction pathways and thus contribute to the transformation of the nasopharyngeal epithelium [611].

**Cytogenetics and comparative genomic hybridization (CGH)**

Only few well-characterised karyotypes of NPC have been described. Despite the many complex rearrangements found, rearrangement and deletion on chromosome 3 have been consistently noted in this cancer [1141, 2707, 2813]. The spectral karyotyping (SKY) analyses have defined the common chromosomal regions of loss including 3p12-p21, 11q14-qter as well as the common regions of gain including 7p15-p14, 7q11.2-q21, 8q21.1-q22, 12q22-q24.1 and 20q were frequently detected [2813]. CGH studies have identified multiple recurrent chromosomal aberrations including loss on chromosomes 3p, 9p, 9q, 11q, 13q, 14q and 16q and gains of 1q, 3q, 12p, and 12q. Common regions of loss are 3p14-21, 14q24-qter, 11q21-qter while common regions of gains are 3q21-26 and 12q13-15 [413, 716, 1148]. Array-based CGH analyses and fluorescence in-situ hybridization (FISH) analyses have identified a cryptic amplification at 3q26 [964, 1149].

**Molecular genetic alterations**

In concordance with CGH results, loss of heterozygosity (LOH) studies have revealed high frequencies of deletion on chromosomes 3p, 9p, 9q, 11q, 13q, 14q and 16q. Multiple minimally deleted regions are identified at 3p14–24.2, 11q21–23, 13q12–14, 13q31–32, 14q24–32, and 16q22–23 [1545]. The characteristic LOH on 3p, 9p, and 14q in almost all tumours suggests that the putative tumour suppressor genes located in these regions probably play important roles in the genesis of NPC. Moreover, deletions on 3p and 9p have been shown to be early events in NPC tumorigenesis [360, 361]. Inactivation of the P16 tumour suppressor gene on 9p21 by homozygous deletion and methylation has been shown to be the most common molecular alteration in NPC tumourigenesis [1541, 1543]. Loss of P16 may result in cell cycle deregulation, while aberrations of the two major cell cycle regulators, P53 and RB, are rare [2453, 2504]. Some studies have also found a high frequency of promoter hypermethylation of RASSF1A, a tumour suppressor gene on 3p21.3, in 70-80% of all cases of primary tumours [1403, 1544]. The tumour suppressor function of RASSF1A may involve the DNA repair system and the RAS-dependent growth control. Other NPC-associated genes in the minimally deleted regions include TSLC1 at 11q23, EDNRB at 13q22, E-CADHERIN and RB2/130 at 16q [457, 1542, 1546, 2642]. Epigenetic inactivation of multiple cancer-associated genes is common in NPC. Aside from P16 and RASSF1A, high frequencies of aberrant methylation are detected in EDNRB (90.5%), RARB2 (80%), DAP-kinase (76%), RIZ1 (60%) and E-CADHERIN (52%) [390, 1403, 1546]. Widespread hypermethylation of CpG islands over the genome imply a “methylator” phenotype in this cancer.

**Expression profiles / Proteomics**

In NPC, P53 mutation is rare, but DNA-P63, a P53 homologue, is consistently over-expressed and may block P53-mediated transactivation and apoptotic network in cancer cells [506, 2453]. Frequent aberrant expression of the cyclin D1, P27 and BCL-2 may also be involved in dysregulation of cell proliferation and apoptosis pathway [100, 1415, 1566]. Overexpression of the hypoxia associated proteins, HIF-alpha, CA IX, and VEGF, is common and associated with poor prognosis [1150]. High MET protein expression level correlates with poor survival in late-stage NPC [2094].

**Genetic susceptibility**

There is strong evidence that genetic predisposition is involved in the genesis of NPC. Epidemiological studies strongly support the existence of susceptible populations in the world: the prevalence of NPC is highly variable in different ethnic groups [1979]; migrants from high-risk areas continue to exhibit high risk of NPC [952, 1501], familial clustering of NPC is frequently observed [2893].

**HLA**

There is an association between HLA phenotype and NPC risk. The association between HLA-A2 and NPC was first reported among Chinese in Singapore [2364]. Subsequent studies have confirmed the association of HLA A2-B46 haplotype with NPC in many different countries [382, 1089, 1223, 1565, 1567, 2365, 2366]. In addition, increased risk of NPC has been found in individuals harbouring HLA B17 in southern China [1567, 2874], Singapore [382] and Malaysia [381]. Haplotypes A2-B17 [2895], A2-B38 {1565}, and A2-B16 [2827] are also shown to be associated with increased risk of NPC. These findings are further supported by linkage or association studies that provide evidence for a NPC predisposing gene in close linkage with the HLA locus [1565, 1567, 1947]. There is a negative association with NPC risk for alleles A11, B13.
and B22 [916]. On the other hand, an association between HLA and NPC has not been found in NPC patients in Alaskan Eskimos, Indians [1427], North Africans [208,1071,1762] and Caucasians [313,314,1769].

**GSTM1 and CYP2E1**
Polymorphism of some metabolic enzyme genes has been reported to influence susceptibility to NPC. Glutathione S-transferase M1 (GSTM1) detoxifies benzopyrene and other carcinogens in tobacco smoke. Studies on association between absence of GSTM1 and increased risk for NPC are conflicting [415,1857]. The cytochrome P450 2E1 (CYP2E1) enzyme catalyzes the metabolic activation of low-molecular weight nitrosamines such as those detected in NPC-associated foods. A variant form of the gene that is detectable by Rsa I digestion (the c2 allele) has been shown to exhibit higher enzymatic activity. If dietary nitrosamines from preserved foods indeed play a direct role in NPC development, exposed individuals possessing different CYP2E1 genotypes may experience differential levels of NPC risk. In a population-based case-control study from Taiwan, individuals possessing the c2/c2 genotype experienced a 2.6-fold risk relative to those with one or two copies of the wild-type allele [1088]. This finding adds to the evidence that nitroamine-containing preserved foods are important nasopharyngeal carcinogens.

**PIGR and TCR**
The mechanism of the entry of EBV into the nasopharyngeal epithelium has not yet been conclusively elucidated, but a receptor on nasopharyngeal epithelial cells, namely polymeric immunoglobulin receptor (PIGR), has been proposed to be involved. It has been reported that one single nucleotide polymorphism (SNP) (1739C->T), located on exon 7 of the gene, is significantly associated with increased risk of NPC [1102]. The SNP is a missense mutation altering the amino acid alanine to valine, and it occurs adjacent to the endoproteolytic cleavage site of the PIGR extracellular domain. It is hypothesized that the homozygous 1739C state may result in the altered efficiency to release IgA-EBV complex and hence increase the possibility of nasopharyngeal epithelial cells to be infected by EBV.

Since T cell receptor (TCR) may mediate immunity against EBV infection, effort has been made to test the association between polymorphism of TCR and NPC. A study has shown NPC susceptibility to be associated dominantly with a 20-kb fragment (P=0.02, RR=8.2) [412].

**Chromosome 4p**
With construction of a human genome genetic linkage map and development of methods and algorithms, linkage analysis has become the robust tool to connect phenotypes with genotypes. A whole genome scan for linkage with NPC has been performed on 32 high risk NPC Cantonese pedigrees [729]. The marker D4S405 on chromosome 4p12–p15 yielded a maximum multipoint lod score of 3.06, a heterogeneity adjusted lod score (HLOD) of 3.21, and a non-parametric linkage score of 2.75 (P=0.005),

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**Fig. 2.20** Nasopharyngeal carcinoma. Spectral karyotyping (SKY) analysis of a nasopharyngeal carcinoma cell line C66-1 exhibiting multiple structural rearrangements.

**Fig. 2.20A** Multistep evolution of nasopharyngeal cancer.
suggesting that a disease susceptibility gene may be linked with D4S405. Fine mapping and haplotype analysis has localized the NPC predisposing gene to chromosome region 4p15.1–q12.

In summary, NPC development may involve susceptibility gene mutations (major genes) and gene polymorphisms (minor-effect genes). In some familial cases, inherited genetic alterations (major gene transmission) could be the first “hit”, and EBV infection may contribute to the second “hit”. Therefore, familial cases usually have a much younger age of onset. However, some other familial cases and probably most sporadic cases may get the first “hit” from both inherited genetic alterations (minor-effect genes, such as HLA, CYP2E1) and somatic genetic changes. In the high prevalence areas like south China, most of the NPC cases belong to this type and they usually have older age of onset than the familial cases with a major gene transmission [2890].

**Prognosis and predictive factors**

The mainstay of treatment for NPC is radiation therapy. Progressive improvement of treatment results for NPC has been reported both from endemic and non-endemic areas. The average 5-year survival steadily increased from around 35% for patients treated in the 1940–60 [1755,1785,2096], to 55-60% in the 1970–90s [1061,1592,2096,2693]. A recent study of patients without distant metastases treated during 1996–2000 showed that a 5-year disease-specific survival (DSS) of 81% and overall survival 75% can now be achieved [1448].

The presenting stage is the most important prognostic factor. A recent study using the 2002 TNM staging System shows that the 5-year DSS for Stage I is 98%, Stage II A-B 95%, Stage III 86%, and Stage IVA-B 73%. In addition, tumour volume may prove to be useful for predicting local control [447,2520].

The importance of host factors varies among different series. In general, younger age (less than 40 years) and female gender are associated with better prognosis [2574]. Interestingly, the influence of age is mainly on local failure, while that of gender is on distant failure. The values of EBV antibodies for predicting prognosis and monitoring disease progression are rather limited [567,2855]. High baseline titers often persist even in patients in remission. Although rising titers to VCA, EA and Zta are associated with disease relapse, the elevation is often not consistently high or early enough for disease monitoring.

Circulating plasma/serum EBV DNA is a more promising prognostic factor. High plasma/serum EBV DNA titers are associated with advanced stages [1549]; both pre-treatment and post-treatment titers correlate significantly with survival [362,1547]. The titer is substantially elevated in patients with active disease (especially distant metastasis), and drops to very low titers upon remission [362,1548,1882,2346].

Aneuploid status or high pre-treatment tumour proliferative fractions, as determined by DNA flow cytometry, correlate significantly with poor survival [2854]. Other biological factors that might have prognostic significance include tumour angiogenesis, c-erbB2 [2209], p53 [1658], nm23-H1 [965], interleukin-10 [828], and vascular endothelial growth factor [2095].

Treatment factors affect the ultimate survival. Significant improvements in treatment results have been attributed to refinement of radiotherapy technique [1454], dose escalation [2561,2576], accelerated fractionation [1447,2717], addition of chemotherapy (concurrent + sequential) [29,1515,2180], and combination of new strategies [2803].

**Fig. 2.22** A putative model for the development of the three forms of nasopharyngeal carcinoma.

**Fig. 2.21** Nasopharyngeal carcinoma. Actuarial disease-specific survival for different stages of nasopharyngeal carcinoma (Source of data: Hong Kong Nasopharyngeal Cancer Study Group on 2687 patients staged with the UICC/AJCC 5th Edition at public centres in Hong Kong during 1996-2000).
Nasopharyngeal papillary adenocarcinoma and salivary gland-type carcinomas

Nasopharyngeal papillary adenocarcinoma

Definition
A low-grade adenocarcinoma characterized by an exophytic growth comprising papillary fronds and glandular structures.

ICD-O code 8260/3

Epidemiology
Nasopharyngeal papillary adenocarcinoma is extremely rare [1902,2672,2770]. The reported age range is 11-64 years (median 37 years) [1902,2672,2770]. Gender distribution is nearly equal [1902,2672,2770].

Localization
The tumour most commonly involves the roof, lateral wall and posterior wall of the nasopharynx [2770].

Clinical features
Nasal obstruction is the main presenting symptom. The diagnosis can be readily confirmed by endoscopic biopsy.

Macroscopy
The tumours are soft or gritty and exophytic, with a papillary, polypoid, or cauliflower appearance. The tumours measure up to 4 cm (median size 2.5 cm) [2770].

Tumour spread and staging
The tumours usually remain confined within the nasopharynx except one reported case with extensive local invasion [1902].

Histopathology
Nasopharyngeal papillary adenocarcinoma arises from the surface epithelium [2770]. The tumour comprises arborizing delicate papillary fronds and crowded glands. The lining columnar or pseud Stratified cells have bland, round to oval nuclei and tiny nucleoli. Mitotic figures are rare; necrosis may be focally identified. Psammoma bodies may be found in some cases. The tumours are unencap- sulated and infiltrate into the surrounding stroma. Diastase-resistant, periodic acid-Schiff intracytoplasmic positive material is present; intraluminal and intracytoplasmic mucicarmine staining may be focally identified. Immunohistochemical staining shows positive reactivity for epithelial markers (i.e., cytokeratin, epithelial membrane antigen), but there is no reactivity for thyroglobulin and S-100 protein. There is no association with Epstein-Barr virus.

Genetic susceptibility
A case has been reported in a patient with Turner syndrome [1902].

Prognosis and predictive factors
This is an indolent low-grade malignant neoplasm with no metastatic potential. It has an excellent prognosis if a complete excision can be achieved [2770].

Salivary gland-type carcinomas

These are very rare in the nasopharynx [2448]. Men are affected nearly three times more frequently than women [1389]. The age range is from 15-74 years with a median age of 50 years. The most frequent types are, in order of frequency, adenoid cystic carcinoma, mucoepidermoid carcinoma and adenocarcinoma not otherwise specified [2273]. Carcinomas at this site frequently present at an advanced stage and often with invasion of the base of the skull, intracranial extension and involvement of the cranial nerves. Adenoid cystic carcinomas [336,1449,2273,2718] are typically insidious in onset, and symptoms may include middle ear effusion, epistaxis, diplopia and symptoms due to cranial nerve palsy (such as pain, paraesthesia, anaesthesia). The microscopic features are similar to those of adenoid cystic carcinoma elsewhere. The 5 and 10 year survival are 78% and 49.5% respectively, and 35% of patients will develop metastasis to bone or lung [2718]. Mucoepidermoid carcinomas [1321,1389,2273] are microscopically similar to those in other sites but rarely psammoma bodies can be seen. Other rare salivary gland-type carcinomas of the nasopharynx include epithelial-myoepithelial carcinoma [1174], myoepithelial carcinoma [1899], acinic cell carcinoma [1890] and polymorphous low-grade adenocarcinoma [1469,2763].

Fig. 2.23 A Nasopharyngeal papillary adenocarcinoma. The tumour comprises complex papillae and glands lined by columnar to spindly cells with bland-looking nuclei. B Mucoepidermoid carcinoma of nasopharynx. There are solid islands of squamoid cells and clear cells.
Benign epithelial tumours

**Hairy polyp**

**Definition**
A presumed developmental anomaly that clinically manifests as a polyp covered by skin with hair and sebaceous glands.

**Synonyms**
Teratoid polyp, dermoid polyp.

**Epidemiology**
Hairy polyps occur in newborns and older infants. There is an unexplained female predominance (female to male ratio 6:1) [1296].

**Localization**
The lateral wall of the nasopharynx, the superior nasopharyngeal aspect of the soft palate, and the tonsils are classic locations for hairy polyps [1296]. They also have been infrequently reported in the middle ear [1310]. No cases have been reported in the sinonasal tract.

**Clinical features**
The usual clinical presentation is a pedunculated mass in the oropharynx or nasopharynx of a newborn or older infant. In the middle ear, the hairy polyps cause recurrent otitis media that is not responsive to the usual treatment. There are individual reports of associated cleft palate or multiple congenital anomalies, including the Dandy-Walker malformation [88].

**Histopathology**
The surface of the polyp is composed of skin with a delicate hyperkeratotic layer and pilosebaceous units. The core is formed by fibroadipose tissue often with foci of cartilage, muscle and bone. Hairy polyps are distinguished from teratomas by a lack of endodermal components.

**Histogenesis**
It has been argued that these polyps are congenital anomalies of the first branchial cleft or choristomas [1045].

**Prognosis and predictive factors**
Complete surgical excision is curative.

**Schneiderian-type papilloma**

**Definition**
A benign tumour that arises from the surface epithelium of the nasopharynx and resembles Schneiderian papillomas of the sinonasal tract [81,1924].

**ICD-O code** 8121/0

**Synonyms**
Fungiform papilloma, inverted papilloma, transitional papilloma, nasopharyngeal papilloma.

**Epidemiology**
Schneiderian-type papillomas of the nasopharynx are distinctly uncommon. They occur in older individuals (mean 62 years, range 45-79) and are 2-3 times more common in males [2499].

**Etiology**
Anatomically, the posterior choanae represent the boundary between the ectodermally-derived (Schneiderian membrane) and endodermally-derived respiratory mucosa that, respectively, line the sinonasal tract and nasopharynx. It is thought that aberrant embryologic displacement of normal Schneiderian mucosa might account for these lesions in the nasopharynx.

**Clinical features**
Most do not exceed two cm in greatest dimension. They are often incidental findings or, at most, result in nasal airway obstruction. The more common Schneiderian papilloma of the sinonasal tract with secondary involvement of the nasopharynx must be excluded before accepting the lesion as primary in the nasopharynx.

**Histopathology**
They are similar to those occurring in the nasal cavity and paranasal sinuses (see section on sinonasal papillomas). Most are of the inverted type (ICD-O code 8121/1).

**Prognosis and predictive factors**
Transnasal or transoral excision is the treatment of choice. Local recurrences are not uncommon. At least one case has been associated with a separate focus of nasopharyngeal squamous cell carcinoma [2499].

**Squamous papilloma**
Squamous papillomas are uncommon in the nasopharynx, and they are morphologically similar to those found in the larynx. See chapter on ‘Tumours of the hypopharynx, larynx and trachea’.

**Ectopic pituitary adenoma**

**Definition**
A benign pituitary gland neoplasm occurring separately from, and without involvement of the sella turcica (i.e., with normal anterior pituitary gland).

**Prognosis and predictive factors**
Ectopic pituitary adenoma of the nasopharynx appearing as a submucosal and unencapsulated cellular tumour; the nasopharyngeal surface epithelium is intact and seen on top.
Tumours of the nasopharynx

ICD-O code 8272/0

Synonyms
Extrasellar pituitary adenoma; extrasellar adenohypophysial tissue; extracranial pituitary adenoma; sphenoidal pituitary adenoma; adenomatous pharyngeal pituitary.

Epidemiology
Ectopic pituitary adenomas of the upper aerodigestive tract are rare, and predominantly occur in adults but have been identified over a wide range of ages, from 16 – 84 years, with a reported mean and median age at presentation of 49 years and 58 years respectively {1425,2752}. Females are affected more often than men {1425,2752}.

Etiology and pathogenesis
The etiology of ectopic pituitary adenomas is unknown. Extrasellar involvement by a pituitary adenoma can result from downward extension of a sellar-based pituitary tumour or occur as an adenomatous tumour arising from ectopic pituitary tissues. The latter may occur from two sources, including embryologic rests along the course of the cephalic invagination of Rathke’s pouch (infrasellar) or anterior pituitary cells attached to the supradiaphragmatic portion of the pituitary stalk. This discussion will be limited to infrasellar-derived ectopic pituitary adenomas.

Localization
In the upper aerodigestive tract, ectopic pituitary adenoma most commonly occurs in the sphenoid bone and sinus {55,1425,1538,2627} and nasopharynx {417,488,2752}. Other sites of occurrence include the nasal cavity {1131,2129,2752}, ethmoid sinus {2752}, and temporal bone {2129}.

Clinical features
The clinical presentation of ectopic pituitary adenomas is primarily related to its space-occupying effects, and include airway obstruction, chronic sinusitis, headache, epistaxis, cerebrospinal fluid leakage and visual field defects {237,417,1425,1667,2627,2752}. Clinical evidence of hormonally active tumours can be identified in over half of the cases {1425} and include Cushing disease {235,309,1261,1538,2274,2396}, acromegaly {491,685,2725} hyperparathyroidism {1538}, hyperthyroidism {488}, amenorrhea {1051}, and hirsutism {2752}. Radiographic imaging is helpful in localizing the lesion and determining the relationship to the sella turcica {2396}. Of note is the fact that the extent of tumour and erosion of bone do not completely correlate with severity of clinical signs and symptoms {1425}.

Furthermore, erosion of the sella turcica does not exclude an ectopic origin {1425}.

Macroscopy
Grossly, it is a polypoid and pedunculated mass ranging in size from 0.7-7.5 cm {1538}. It is usually solitary; rare examples may occur synchronously in separate sites {237}.

Histopathology
Ectopic pituitary adenomas are submucosal and unencapsulated cellular tumours with solid, organoid and trabecular growth patterns; tumour nests are separated by fibrovascular stroma. The neoplastic cells have round to oval nuclei with dispersed nuclear chromatin, inconspicuous to small nucleoli, and granular cytoplasm that can be eosinophilic, amphophilic or clear. Extracellular stromal hyalinization may be prominent. Nuclear pleomorphism, mitotic activity and necrosis are uncommon. The surface epithelium is intact and unremarkable. The tumour effaces the normal structures in the submucosa although residual seromucous glands can be identified.

Immunoprofile
Ectopic pituitary adenomas show strong cytoplasmic immunoreactivity for cytokeratin, synaptophysin, chromogranin, neuron specific enolase, and may stain for a variety of pituitary hormones including adrenocorticotrophic hormone (ACTH), prolactin, thyroid stimulating hormone (TSH), follicle stimulating hormone (FSH), growth hormone (GH), and luteinizing hormone (LH). Tumours may demonstrate immunoreactivity with only a single pituitary hormone (monohormonal pituitary adenoma), multiple hormones (plurihormonal pituitary adenoma) or no pituitary hormone (null cell pituitary adenoma).

Electron microscopy
Intracytoplasmic secretory granules of varying numbers can be identified.

Prognosis and predictive factors
Surgical resection is the treatment of choice for smaller, accessible tumours {1425}; complete surgical eradication is usually curative {2752}. However, complete surgical resection may not be possible for larger, invasive tumours. When resection is incomplete or cannot be accomplished due to size and extent of

Fig. 2.25 Ectopic pituitary adenoma. A Organoid growth pattern. B Growth in the form of ribbons. C Scattered mitotic figures are seen in a tumour that otherwise shows uniform neoplastic cells with minimal pleomorphism and dispersed chromatin. D Extension into bone may be present.
the tumour, postoperative radiotherapy is indicated [55,1425,2627,2666]. Dopamine agonist drugs (e.g., bromocriptine) have been effective in reducing the size (not permanently) in prolactin-secreting adenomas and the mitotic rate of other pituitary adenomas [1425]. Somatostatin analog treatment with octreotide has been shown to reduce tumour size and may, in the proper setting of a pituitary adenoma with high somatostatin receptor content, be administered in lieu of surgery in patients whose tumours are too large to be adequately resected [2093]. A rare example of malignant transformation of an ectopic pituitary adenoma has been reported [1131].

Salivary gland anlage tumour

Definition
A benign tumour with mixed epithelial and mesenchymal elements, recapitulating the early stages in the embryology of the salivary gland between the 4th and 8th weeks of development.

Synonym
Congenital pleomorphic adenoma

Epidemiology
Fewer than 20 cases have been reported in the literature [229,233,296,572,1007, 1168,1720,1763]. Most patients are diagnosed in the immediate neonatal period or by the age of 6-weeks. Males exceed females by a 13:3 ratio.

Clinical features
Almost all patients present with respiratory and feeding difficulties. Bleeding has rarely been reported. Clinical examination reveals a midline pedunculated erythematous polyp [229].

Macroscopy
A firm, smooth to lobulated mass measuring between 1.3 and 3 cm in greatest dimension is the typical gross appearance. The surface is usually glistening. The remnants of a stalk may or may not be apparent. A vague nodularity is appreciated on its greyish-tan to reddish cut surface. Cysts and interstitial haemorrhage may occur.

Histopathology
A non-keratinizing squamous mucosa overlies multiple contiguous cellular nodules. The nodules are separated by fibrous and myxoid stroma containing duct-like structures and nests of solid or cystic squamous epithelium. In areas, the duct-like structures are connected to the surface epithelium. The epithelial units within the internodular stroma blend into the cellular nodules, which are comprised of fusiform cells forming short fascicles or trabecular structures, interspersed with poorly formed tubules with or without lumens. The fusiform cells have eosinophilic cytoplasm with indistinct cell borders. The nuclei are bland and uniform, and mitotic activity is quite low. The interstitium can show haemorrhage, and rarely bone formation [296].

Immunoprofile
The cellular nodules display a mixed pattern of reactivity for vimentin, cytokeratin and actin and are generally non-reactive for S-100 protein and glial fibrillary acidic protein. Nascent tubules and ducts within the stromal nodules show a luminal pattern of positivity for epithelial membrane antigen. The differentiated epithelial components are reactive for pancytokeratin and cytokeratin 7; epithelial membrane antigen positivity is restricted to the tubular structures. Salivary gland amylase is expressed consistently.

Prognosis and predictive factors
Complete excision is curative in virtually all cases.

Craniopharyngioma

ICD-O code
9350/1

Exceptionally, craniopharyngioma can arise in the nasopharynx or involves the nasopharynx through downward invasion from a suprasellar location. The morphological features are identical to the suprasellar counterpart [316,1609,1612, 2062,2083,2525].
The spectrum and clinicopathological features of nasopharyngeal soft tissue tumours are similar to those of other sites in the upper aerodigestive tract, except for angiofibroma, which typically presents in the nasopharynx.

**Definition**
A benign, highly cellular and richly vascularized mesenchymal neoplasm that involves the nasopharynx in males.

**ICD-O code**
9160/0

**Synonyms**
Juvenile nasopharyngeal angiofibroma; angiofibroma; fibroangioma; fibroma

**Epidemiology**
Nasopharyngeal angiofibroma represents <1% of all nasopharyngeal tumours [190,267,512,1434,1503,1861,2654]. Boys and adolescent to young men are almost exclusively affected, with a peak in the 2nd decade of life. If a female is affected, testicular feminisation has to be excluded. Fair-skinned and red-haired males are more commonly affected.

**Etiology**
There is no known etiology although testosterone-dependent puberty-induced tumour growth may be ameliorated by blockade of estrogen or progesterone receptors within the tumour [717,1861].

**Localization**
This tumour arises in the posterolateral nasal wall or the nasopharynx. There is often extensive infiltration into the surrounding tissues [190,267,512,1434,1503,1861,2654].

**Clinical features**
Patients usually present with nasal obstruction and/or recurrent, spontaneous epistaxis, nasal discharge, facial deformity (including proptosis), diplopia, exophthalmos, sinusitis, otitis media, tinnitus, rhinolalia, deafness, headaches, dyspnoea, and rarely, anosmia or pain [190,267,512,1434,1503,1861,2654].

**Imaging**
Routine radiographs reveal a soft tissue density in the nasopharynx in conjunction with anterior bowing of the posterior wall of the maxillary sinus as well as distortion and posterior displacement of the pterygoid plates (Holman-Miller sign). Bony margins may be eroded, but are distinct. Computed tomography allows for accurate determination of the extent of the disease as well as the best possible surgical approach. Angiography allows for identification of the feeding vessel(s) and pre-surgical embolization. Tumour blush on angiogram is characteristic [1434,2654]. Due precautions have to be taken in obtaining biopsies from the lesion because of the risk of life-threatening bleeding.

**Macroscopy**
The tumours range in size up to 22 cm, with a mean of about 4 cm. They are polypoid with a rounded or multinodular contour, with red, grey-tan cut surfaces [190,267,512,1434,1503,1861,2654].

**Tumour spread and staging**
The tumour expands in all directions from the nasopharyngeal region, following the path of least resistance: anteriorly into the nasal cavity and maxillary sinuses, laterally into the pterygoid region, temporal fossa and infratemporal fossa (resulting in a cheek or intraoral buccal mass); superiorly into orbit and middle cranial fossa; or to the opposite side. This type of extensive involvement is seen in up to 30% of cases, explaining the potential aggressive nature of this benign neoplasm [190,267,512,1434,1503,1861,2654].

A number of staging systems have been suggested, [384,767,2111,2309] with a modification based on size and location used most frequently.

**Histopathology**
There is a vascular proliferation set in a fibrous stroma. The vessels are mostly thin-walled, slit-like (“staghorn”) or dilated with calibres ranging from capillary size to large, patulous vessels. The mus-
Nasopharyngeal angiofibroma

The vascular layer can be absent, focal and pad-like, or circumferential. Endothelial cells may be plump but are usually attenuated. The fibrous stroma consists of plump spindle, round, angular, or stellate shaped cells and a varying amount of fine and coarse collagen fibres; background myxoid degeneration is common (especially in embolized specimens). The nuclei of the stromal cells are generally cytologically bland, but they may be multinucleated or show some degree of pleomorphism in the more cellular areas. Mast cells may be seen, but other inflammatory elements are usually absent (except when there is surface ulceration) (190,267,512,1434,1503,1861,2654). Long-standing lesions show increased fibrosis and diminished vasculature. Treatment with hormones results in increased collagenization of the stroma with fewer, but thicker-walled vessels. In specimens excised after embolization, the tumour often shows areas of infarction, and emboli can be seen in some blood vessels. Sarcomatous transformation is an exceedingly uncommon event, usually following radiation therapy (2431).

Immunoprofile

Occasional elastic fibres can be identified in the vessel walls, although they are generally absent in the stroma. The vessel wall cells are immunoreactive with vimentin and smooth muscle actin (SMA), whereas the stromal cells are immunoreactive with vimentin only, except in areas of increased fibrosis, where focal SMA may be identified. Desmin may be focally immunoreactive in larger vessels at the periphery of the tumour. Stromal and endothelial cells are variably reactive with androgen and estrogen/progesterone receptors. Factor VIII R-Ag, CD34 and CD31 highlight the endothelium, but not the stromal cells. The stromal cells are negative for S-100 protein (190,1503). Platelet derived growth factor B and insulin-like growth factor type II are both over-expressed (1812).

Electron microscopy

Ultrastructurally, the stromal cells contain lobulated nuclei, intranuclear inclusions, variable amounts of rough endoplasmic reticulum and thin filaments, hemidesmosomes, focal basal lamina and prominent pinocytotic vesicles, suggesting a hybrid mesenchymal cell (myofibroblast) (2565).

Differential diagnosis

The differential diagnosis includes lobular capillary haemangioma (pyogenic granuloma), nasal inflammatory polyps with fibrosis or atypical stromal cells, antrochoanal polyps, and peripheral nerve sheath tumour.

Histogenesis

It has been proposed that the tumour arises from a fibrovascular nidus that lies dormant until puberty, when testosterone stimulates tumour growth (1861).

Genetic susceptibility

There are isolated reports of an association with familial adenomatous polyposis (757,885).

Prognosis and predictive factors

This benign tumour is characterized by local aggressive growth, with recurrences in about 20% of patients (>50% in older series), most commonly intracranially, and usually within the first 2 years after diagnosis. Patients may be managed with selective angiographic embolization or hormonal therapy prior to definitive surgical resection. Radiation therapy has been successfully implemented to manage large, intracranial, or recurrent tumours, but surgery is still the therapy of choice (190,267,512,1434,1503,1861,2654).

### Table 2.05 System for staging nasopharyngeal angiofibroma (384,767,2309).

<table>
<thead>
<tr>
<th>Stage</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Tumour limited to the nasopharynx with no bone destruction</td>
</tr>
<tr>
<td>II</td>
<td>Tumour invading the nasal cavity, maxillary, ethmoid, or sphenoid sinuses with no bone destruction</td>
</tr>
<tr>
<td>III</td>
<td>Tumour invading the pterygopalatine fossa, infra-temporal fossa, orbit or parasellar region</td>
</tr>
<tr>
<td>IV</td>
<td>Tumour with massive invasion of the cranial cavity, cavernous sinus, optic chiasm, or pituitary fossa</td>
</tr>
</tbody>
</table>

### Fig. 2.28 Nasopharyngeal angiofibroma.

**A** Thin walled vessels surrounded by dense, “keloid-like” collagen. Stellate fibroblasts are noted. **B** Smooth muscle-walled vessels, patulous vessels and capillaries are all surrounded by the characteristic collagenized stroma. **C** A large thin-walled vessel is associated with fibrous connective tissue, inflammatory cells and stellate fibroblasts. **D** Heavily collagenized stroma demonstrates only a few stellate fibroblastic cells.
**Hodgkin lymphoma**

Hodgkin lymphoma only rarely shows primary involvement of the nasopharynx \{1274,1602,1756,1763,1922\}. The patients usually present with nasal obstruction or otitis media, and frequently have low stage (stage I/II) disease. Most of the tumours are of mixed cellularity and nodular sclerosis subtypes. The majority of cases involving the nasopharynx are associated with Epstein-Barr virus \{1274,1756\}. Please refer to ‘Hodgkin lymphoma’ in ‘WHO classification of tumours: Tumours of haematopoietic and lymphoid tissues’ for details.

**Non-Hodgkin lymphoma**

**Definition**

Primary non-Hodgkin lymphoma (NHL) of the nasopharynx is defined as a lymphoid cell neoplasm in which the bulk of disease occurs in this site.

**Epidemiology**

Nasopharyngeal NHL accounts for 2.5% of all extranodal NHLs \{809\}. Most cases have been reported in the literature in combination with either NHL of the nasal cavity or NHL of the Waldeyer ring, rendering it difficult to extract the specific details on nasopharyngeal NHL \{420,1704,2250,2849\}. In some cases, there is simultaneous involvement of both the nasopharynx and nasal cavity, precluding determination of the site of origin of the NHL.

In the West, nearly all cases of nasopharyngeal NHL are of B-cell lineage (most commonly diffuse large B-cell lymphoma, DLBCL) \{1704\}. The situation is different in Asia, where B-cell lymphomas account for only 50-60% of cases \{420,2849\}, due to a higher frequency of extranodal NK/T cell lymphomas and peripheral T-cell lymphomas.

Most patients with nasopharyngeal NHL are adults. Patients with extranodal NK/T cell lymphoma of nasal-type have a male to female ratio of 3:1, and a median age of 53 years \{420\}. Patients with B-cell lymphomas are generally one decade older (median age of 63 years), and the male to female ratio is only 1.2:1 \{420\}. Burkitt lymphoma occurs more frequently in children and young adults \{2826\}.

**Etiology**

The etiology is unknown, except that extranodal NK/T cell lymphoma of nasal-type is strongly associated with Epstein-Barr virus (EBV) (>95%) irrespective of the ethnic background of the patients \{1195\}. The association of nasopharyngeal DLBCL with EBV is weak \{376\}.

**Clinical features**

The patients present with nasal obstruction, epistaxis, hearing impairment, dysphagia, headache or neck mass, similar to the presenting symptoms of nasopharyngeal carcinoma. A small proportion of patients have concurrent cervical lymphadenopathy, a feature seen more frequently in DLBCL than extranodal NK/T cell lymphoma.

**Tumour spread and staging**

The majority (80%) of patients have localized disease (Stage I/E/IIE) at presentation \{420,1500,1505,1550\}. Extranodal NK/T cell lymphoma tends to disseminate to various sites, such as skin, gastrointestinal tract, liver, lymph node and testis, during the course of disease. There is a propensity for DLBCL to spread to the cervical lymph nodes \{420\}.

**Histopathology**

DLBCL and extranodal NK/T cell lymphoma of nasal-type occurring in the nasopharynx are morphologically and morphologically and morphologically...
immunophenotypically similar to those seen in the nasal cavity. Other types of NHL, for example, Burkitt lymphoma, follicular lymphoma, mantle cell lymphoma, extranodal marginal zone B-cell lymphoma of MALT type, and peripheral T-cell lymphoma unspecified may also affect the nasopharynx, but at a much lower frequency [420,1704,2849]. Please refer to the sections of ‘non-Hodgkin lymphoma’ in ‘Tumours of the nasal cavity and paranasal sinuses’ and ‘Tumours of the oral cavity and oropharynx’ for details.

**Differential diagnosis**

Distinction between nasopharyngeal carcinoma and DLBCL can be difficult at times because the carcinoma cells in nasopharyngeal carcinoma can appear discohesive due to submergence in a dense lymphoplasmacytic infiltrate, while DLBCL can sometimes form tight cell clusters. Positive immunostaining for cytokeratin would support the former diagnosis, and expression of lymphoid markers (including CD20) would support the latter. Infectious mononucleosis involving the nasopharynx can also mimic DLBCL [2547], but can be suspected or recognized by the young age of the patient, presence of a range of large cells with apparent maturation to plasmablasts and plasma cells, lack of frank cytologic atypia, and polyclonal immunoglobulin staining in the large cells. Extranodal NK/T cell lymphoma with small cell predominance can be difficult to recognize as being a malignant neoplasm. Histologic features suggestive of the diagnosis include extensive effacement of architecture, marked coagulative necrosis, angiocentric growth, and wide separation of the mucosal glands. The diagnosis is supported by the demonstration of sheets of CD56+, CD3ε+, EBER+ cells. In the rare case of herpes simplex infection involving the nasopharynx, there can be a dense lymphoid infiltrate with extensive CD56 expression, causing confusion with extranodal NK/T cell lymphoma. In contrast to NK/T cell lymphoma, these CD56+ cells express CD4 and CD5, and there is no association with EBV. The diagnosis is confirmed by identifying the herpes simplex virus-infected multinucleated giant cells with ground glass nuclei with or without nuclear inclusions, which can be further confirmed by immunostaining for herpes simplex virus [2523].

**Prognosis and predictive factors**

Radiotherapy is the treatment of choice for extranodal NK/T cell lymphoma, often in combination with additional treatment modalities [421]. Chemotherapy and/or radiotherapy is usually given for patients with DLBCL. The overall survival rate for extranodal NK/T cell lymphoma of nasal-type is only 30-50% [421,422,1312,1838]. Factors associated with worse outcome include: advanced stage, poor performance status, B symptoms and bulky disease [422]. B-cell lymphomas show a slightly more favourable outcome [420].

**Follicular dendritic cell sarcoma / tumour**

Follicular dendritic cell sarcoma/tumour is a rare tumour showing morphologic, immunophenotypic and ultrastructural features of follicular dendritic cells. Primary involvement of the nasopharynx is rare [189,359], and may arise from an underlying hyaline-vascular Castleman disease [359]. Please refer to the section of ‘Follicular dendritic cell sarcoma/tumour’ in ‘Tumours of the oral cavity and oropharynx’ for details.
Tumours of bone and cartilage

The spectrum and clinicopathological features of nasopharyngeal tumours of bone and cartilage are similar to those of other sites in the upper aerodigestive tract, except for chordoma, which typically presents in the nasopharynx.

Chordoma

Definition
A low-grade malignant tumour that recapitulates the notochord.

ICD-O code 9370/3

Epidemiology
Chordomas account for approximately 4% of malignant bone tumours [2655]. About a third involve the base of the skull, and a small proportion may involve the nasopharynx and/or paranasal sinuses. There is a male predilection. The patients are predominantly adults, but children can also be affected.

Clinical features
Patients usually present with non-specific symptoms, such as headache, nasal obstruction, and symptoms related to cranial nerve involvement. Rarely, they present with nasal polyps [325]. Imaging studies show lytic destruction of the basisphenoid centred in the clivus. The tumour frequently extends into the middle cranial fossa and nasopharynx. Calcification is occasionally seen.

Histopathology
Chordomas typically show a lobulated growth pattern. Polygonal or ovoid tumour cells are arranged in cords, lobules and sheets in a myxoid background. The nuclei are typically round and uniform, but may exhibit considerable pleomorphism. The cytoplasm is abundant and eosinophilic, and at times clear. Vacuolated cells (physaliferous cells) are present to a variable degree. The tumour cells are immunoreactive for cytokeratins, epithelial membrane antigen and S100 protein.

The main differential diagnoses are epithelial neoplasms (such as mucinous carcinoma, salivary gland tumours, poorly differentiated carcinoma) and chondrosarcoma. The lobulation, physaliferous cells and diffuse strong S100 protein immunoreactivity distinguish chordoma from carcinoma. Chondrosarcoma is negative for cytokeratin.

Prognosis and predictive factors
Chordoma is a low-grade tumour and distant metastases are rare. Chordomas involving the nasopharynx are often treated by radiation therapy because complete surgical resection is practically impossible because of the anatomy [709].

Secondary tumours

Definition
Tumours that metastasize to the nasopharynx from other primary malignancies. Direct invasion from tumours of adjacent sites, leukemias and lymphomas are excluded.

Epidemiology
Metastases to the nasopharynx are extremely rare [1685]. The majority of patients are over the age of 50 years. Reported primary tumours and tumour sites include malignant melanoma (cutaneous) 9 cases, kidney (3 renal cell, 1 Wilms), lung (4 cases), and one case each of breast, colon and cervical cancer [1685].

Clinical features
Patients may be asymptomatic or present with nasal obstruction, epistaxis, unilateral serous otitis media secondary to blockage of the eustachian tube, or otalgia. Large bulky metastases can extend into the nasal cavity or deform the soft palate. A long disease-free interval between treatment of the primary tumour and the appearance of metastasis in the nasopharynx may confuse the diagnosis, raising the possibility of a new primary neoplasm of the nasopharynx. However, this is not uncommon for malignant melanoma and renal cell carcinoma.

Pathogenesis
Most metastases to the nasopharynx are haematogenous, possibly arising in some instances through Batson's paravertebral venous plexus.

Prognosis and predictive factors
Metastasis is an ominous sign associated with a poor prognosis.
Squamous cell carcinoma is, by far, the most important tumour of the hypopharynx, larynx and trachea. It is clearly related to the abuse of tobacco and alcohol and, as such, could be drastically reduced if individuals would only alter their lifestyles. Precursor lesions have been identified and the genetic-molecular events underlying their origin and progression into clinically apparent carcinomas are gradually being elucidated. The terminology of these precursor lesions, however, is still evolving and no single classification has been universally accepted. The three most commonly used classifications and their equivalent terms are presented.
### WHO histological classification of tumours of the hypopharynx, larynx and trachea

#### Malignant epithelial tumours

<table>
<thead>
<tr>
<th>Tumour Type</th>
<th>Morphology Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Squamous cell carcinoma</td>
<td>8070/3</td>
</tr>
<tr>
<td>Verrucous carcinoma</td>
<td>8051/3</td>
</tr>
<tr>
<td>Basaloid squamous cell carcinoma</td>
<td>8083/3</td>
</tr>
<tr>
<td>Papillary squamous cell carcinoma</td>
<td>8052/3</td>
</tr>
<tr>
<td>Spindle cell carcinoma</td>
<td>8074/3</td>
</tr>
<tr>
<td>Acantholytic squamous cell carcinoma</td>
<td>8075/3</td>
</tr>
<tr>
<td>Adenosquamous carcinoma</td>
<td>8560/3</td>
</tr>
<tr>
<td>Lymphoepithelial carcinoma</td>
<td>8082/3</td>
</tr>
<tr>
<td>Giant cell carcinoma</td>
<td>8031/3</td>
</tr>
<tr>
<td>Malignant salivary gland-type tumours</td>
<td></td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td>8430/3</td>
</tr>
<tr>
<td>Adenoid cystic carcinoma</td>
<td>8200/3</td>
</tr>
</tbody>
</table>

#### Neuroendocrine tumours

<table>
<thead>
<tr>
<th>Tumour Type</th>
<th>Morphology Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Typical carcinoid</td>
<td>8240/3</td>
</tr>
<tr>
<td>Atypical carcinoid</td>
<td>8249/3</td>
</tr>
<tr>
<td>Small cell carcinoma, neuroendocrine type</td>
<td>8041/3</td>
</tr>
<tr>
<td>Combined small cell carcinoma, neuroendocrine type</td>
<td>8045/3</td>
</tr>
</tbody>
</table>

#### Benign epithelial tumours

<table>
<thead>
<tr>
<th>Tumour Type</th>
<th>Morphology Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Papilloma</td>
<td>8050/0</td>
</tr>
<tr>
<td>Papillomatosis</td>
<td>8060/0</td>
</tr>
<tr>
<td>Salivary gland-type adenomas</td>
<td></td>
</tr>
<tr>
<td>Pleomorphic adenoma</td>
<td>8940/0</td>
</tr>
<tr>
<td>Oncocytic papillary cystadenoma</td>
<td>8290/0</td>
</tr>
</tbody>
</table>

#### Soft tissue tumours

<table>
<thead>
<tr>
<th>Tumour Type</th>
<th>Morphology Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Malignant tumours</td>
<td></td>
</tr>
<tr>
<td>Fibrosarcoma</td>
<td>8810/3</td>
</tr>
<tr>
<td>Malignant fibrous histiocytoma</td>
<td>8830/3</td>
</tr>
<tr>
<td>Liposarcoma</td>
<td>8850/3</td>
</tr>
<tr>
<td>Leiomyosarcoma</td>
<td>8890/3</td>
</tr>
</tbody>
</table>

#### Rhabdomyosarcoma 8900/3

#### Angiosarcoma 9120/3

#### Kaposi sarcoma 9140/3

#### Malignant peripheral nerve sheath tumour 9040/3

#### Synovial sarcoma 9040/3

#### Borderline tumours / LMP

<table>
<thead>
<tr>
<th>Tumour Type</th>
<th>Morphology Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inflammatory myofibroblastic tumour</td>
<td>8825/1</td>
</tr>
</tbody>
</table>

#### Benign tumours

<table>
<thead>
<tr>
<th>Tumour Type</th>
<th>Morphology Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Schwannoma</td>
<td>9560/0</td>
</tr>
<tr>
<td>Neurofibroma</td>
<td>9540/0</td>
</tr>
<tr>
<td>Lipoma</td>
<td>8850/0</td>
</tr>
<tr>
<td>Leiomyoma</td>
<td>8890/0</td>
</tr>
<tr>
<td>Rhabdomyoma</td>
<td>8900/0</td>
</tr>
<tr>
<td>Hemangioma</td>
<td>9120/0</td>
</tr>
<tr>
<td>Lymphangioma</td>
<td>9170/0</td>
</tr>
<tr>
<td>Granular cell tumour</td>
<td>9580/0</td>
</tr>
</tbody>
</table>

#### Haematolymphoid tumours

<table>
<thead>
<tr>
<th>Tumour Type</th>
<th>Morphology Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tumours of bone and cartilage</td>
<td></td>
</tr>
<tr>
<td>Chondrosarcoma</td>
<td>9220/3</td>
</tr>
<tr>
<td>Osteosarcoma</td>
<td>9180/3</td>
</tr>
<tr>
<td>Chondroma</td>
<td>9220/0</td>
</tr>
<tr>
<td>Giant cell tumour</td>
<td>9250/1</td>
</tr>
<tr>
<td>Mucosal malignant melanoma</td>
<td>8720/3</td>
</tr>
</tbody>
</table>

#### Secondary tumours

### Notes

1. Morphology code of the International Classification of Diseases for Oncology (ICD-O) (821) and the Systematized Nomenclature of Medicine (http://snomed.org). Behaviour is coded /0 for benign tumours, /3 for malignant tumours, and /1 for borderline or uncertain behaviour.
### TNM classification of carcinomas of the larynx

| Subglottis | T1 | Tumour limited to subglottis |
| T2 | Tumour extends to vocal cord(s) with normal or impaired mobility |
| T3 | Tumour limited to larynx with vocal cord fixation |
| T4a | Tumour invades through cricoid or thyroid cartilage and/or invades tissues beyond the larynx, e.g., trachea, soft tissues of neck including deep/extrinsic muscle of tongue (genioglossus, hyoglossus, palatoglossus, and styloglossus), strap muscles, thyroid, oesophagus |
| T4b | Tumour invades prevertebral space, mediastinal structures, or encases carotid artery |

| N – Regional lymph nodes# | NX | Regional lymph nodes cannot be assessed |
| N0 | No regional lymph node metastasis |
| N1 | Metastasis in a single ipsilateral lymph node, 3 cm or less in greatest dimension |
| N2 | Metastasis as specified in N2a, 2b, 2c below |
| N2a | Metastasis in a single ipsilateral lymph node, more than 3 cm but not more than 6 cm in greatest dimension |
| N2b | Metastasis in multiple ipsilateral lymph nodes, none more than 6 cm in greatest dimension |
| N2c | Metastasis in bilateral or contralateral lymph nodes, none more than 6 cm in greatest dimension |
| N3 | Metastasis in a lymph node more than 6 cm in greatest dimension |

Note: Midline nodes are considered ipsilateral nodes.

| M – Distant metastasis | MX | Distant metastasis cannot be assessed |
| M0 | No distant metastasis |
| M1 | Distant metastasis |

| Stage Grouping |
| Stage 0 | Tis | N0 | M0 |
| Stage I | T1 | N0 | M0 |
| Stage II | T2 | N0 | M0 |
| Stage III | T1, T2 | N1 | M0 |
| | T3 | N0, N1 | M0 |
| Stage IVA | T1,T2,T3, | N2 | M0 |
| | T4a | N0, N1, N2 | M0 |
| Stage IVB | T4b | Any N | M0 |
| | Any T | N3 | M0 |
| Stage IVC | Any T | Any N | M1 |

---

1 (847,2418).
2 A help desk for specific questions about the TNM classification is available at www.uicc.org/index.php?id=508
3 # The regional lymph nodes are the cervical nodes.
## TNM classification of carcinomas of the hypopharynx

### TNM classification

**T** – Primary tumour

**TX** Primary tumour cannot be assessed

**T0** No evidence of primary tumour

**Tis** Carcinoma in situ

**T1** Tumour limited to one subsite of hypopharynx and 2 cm or less in greatest dimension

**T2** Tumour invades more than one subsite of hypopharynx or an adjacent site, or measures more than 2 cm but not more than 4 cm in greatest dimension, without fixation of hemilarynx

**T3** Tumour more than 4 cm in greatest dimension, or with fixation of hemilarynx

**T4a** Tumour invades any of the following: thyroid/cricoid cartilage, hyoid bone, thyroid gland, oesophagus, central compartment soft tissue*

**T4b** Tumour invades prevertebral fascia, encases carotid artery, or invades mediastinal structures.

*Note: Central compartment soft tissue includes prelaryngeal strap muscles and subcutaneous fat.

### N – Regional lymph nodes

**NX** Regional lymph nodes cannot be assessed

**N0** No regional lymph node metastasis

**N1** Metastasis in a single ipsilateral lymph node, 3 cm or less in greatest dimension

**N2** Metastasis as specified in N2a, 2b, 2c

**N2a** Metastasis in a single ipsilateral lymph node, more than 3 cm but not more than 6 cm in greatest dimension

**N2b** Metastasis in multiple ipsilateral lymph nodes, none more than 6 cm in greatest dimension

**N2c** Metastasis in bilateral or contralateral lymph nodes, none more than 6 cm in greatest dimension

**N3** Metastasis in a lymph node more than 6 cm in greatest dimension

### Stage Grouping

<table>
<thead>
<tr>
<th>Stage</th>
<th>Tis</th>
<th>N0</th>
<th>M0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage 0</td>
<td>Tis</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage I</td>
<td>T1</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage II</td>
<td>T2</td>
<td>N0</td>
<td>M0</td>
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<tr>
<td>Stage III</td>
<td>T1, T2</td>
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<td>M0</td>
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<tr>
<td></td>
<td>T3</td>
<td>N0, N1</td>
<td>M0</td>
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<td>Stage IVA</td>
<td>T1, T2, T3</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IVB</td>
<td>T4a</td>
<td>N0, N1, N2</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T4b</td>
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<td>M0</td>
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<td>Stage IVC</td>
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<td>Any N</td>
<td>M1</td>
</tr>
</tbody>
</table>

---

1. (947.2418).
2. A help desk for specific questions about the TNM classification is available at www.uicc.org/index.php?id=508
3. ## The regional lymph nodes are the cervical nodes.
With emphasis now on accurate staging and conservative surgery to retain as many functions as possible, especially in the larynx, the pathologist has emerged as an invaluable member of the health care team. Precise and detailed examination of resected head and neck specimens regarding the site of origin of the tumour, structures involved, tumour grade, adequacy of resection margins, and the presence of lymph node metastasis, extranodal spread of tumour, perineural involvement, and vascular invasion are just a few of many features that are important to the clinician who must decide on the total therapeutic regimen for the patient.

Definitions / anatomy

**Larynx**

The larynx extends from the tip of the epiglottis to the inferior border of the cricoid cartilage. Anteriorly, its boundaries are the lingual epiglottis, the thyrohyoid membrane, the anterior commissure, thyroid cartilage, cricothyroid membrane and the anterior arch of the cricoid cartilage. The posterior boundaries include the posterior commissure mucosa (which covers the cricoid cartilage) the arytenoid region, and the interarytenoid space.

The larynx is divided into three compartments - supraglottis, glottis, and subglottis. The supraglottis is composed of the epiglottis, aryepiglottic folds, false vocal cords (vestibular folds), ventricles and saccules. The tip of the epiglottis and the aryepiglottic folds form the superior and lateral supraglottic margins. The inferior limit is a horizontal plane passing through the lateral margin of the ventricle at its junction with the superior surface of the true vocal cord (vocal fold) (947). The ventricle is the “pocket” between the true and false vocal cords. The lateral superior ventricular extension, or “cul-de-sac”, is variably sized, and referred to as the saccule. The epiglottis is further divided into suprathyroid and infrathyroid components by a plane at the level of the hyoid bone.

The glottis extends, superiorly, from a horizontal plane passing through the lateral margin of the ventricle, at its junction with the superior true vocal cord, to an imaginary horizontal plane 10 mm inferiorly from the lateral margin of the ventricle (947). The glottis consists of the true vocal cords, plus their undersurfaces, and the anterior and posterior commissures. The subglottis extends from 10 mm below the true vocal cords to the inferior margin of the cricoid cartilage. Most tumours that clinically appear as “subglottic”, actually arise from the undersurface of the true vocal cord and are still considered glottic. The term “transglottic” does not refer to a specific anatomic site. It designates those tumours that cross the ventricle vertically, to involve both the supraglottis and glottis, and occasionally subglottis (1679). The growth and spread of laryngeal tumours is determined by the site of origin and the anatomic barriers of the different laryngeal compartments (1941). Three of these are especially important: anterior commissure tendon (Broyles’ ligament), paraglottic space and the preepiglottic space.

The anterior commissure tendon is a band of fibrous tissue 1 mm in width and 10 mm in length that extends from the vocal ligaments to the midline of the inner surface of the upper thyroid cartilage (286). It is significant not only because it contains lymphatic and blood vessels, but also because it is devoid of perichondrium at the attachment to the thyroid cartilage, thereby acting as a conduit for tumour spread into the adjacent soft tissue or the prelaryngeal (Delphian) lymph node.

The paraglottic space is a potential space deep to the ventricles and saccules filled with adipose and loose connective tissue. It is bounded by the
conus elasticus inferiorly, the thyroid cartilage laterally, the quadrangular membrane medially, and the pyriform sinus posteriorly. The pre-epiglottic space, also filled with adipose and connective tissue, is triangular shaped. It is bounded anteriorly by the thyroid cartilage and thyrohyoid membrane, posteriorly by the epiglottis and thyroepiglottic ligament, and superiorly by the hyoepiglottic ligament which forms its base. Both paraglottic and pre-epiglottic spaces contain lymphatics and blood vessels, but no lymph nodes. Suprahypopharyngeal epiglottic tumours are distinct from the more common infrapharyngeal tumours in that they are superior to the pre-epiglottic space, and often spread to the base of the tongue. Tumours that invade the pre-epiglottic and paraglottic spaces may spread without impedance through the loose connective tissue and eventually invade the extralaryngeal tissue. The supraglottic larynx is well endowed with lymphatics draining primarily into the upper, middle, and lower jugular lymph nodes (levels II, III and IV, respectively). The glottis, in contrast, has a limited lymphatic supply, but does drain to the same group of lymph nodes. If a glottic carcinoma extends more than one centimeter inferiorly, the paratracheal lymph nodes are at risk for metastasis. The lymphatic drainage of the subglottis is mainly to the paratracheal lymph nodes and, only infrequently, to the middle and lower jugular lymph nodes (levels III and IV). Early in life, the larynx is entirely lined by ciliated respiratory epithelium. With time, this epithelium is gradually replaced by non-keratinizing stratified squamous epithelium. The adult larynx is lined entirely by squamous epithelium, with the exception of the ventricles and the subglottis – which continue to be lined by respiratory epithelium. Infrequently, one may see small patches of persistent ciliated respiratory epithelium in an otherwise typical adult supraglottis. The nonkeratinized squamous mucosa of the true vocal cords is normally about 5–10 cells thick. Although mucoserous glands are abundant in the supraglottis and subglottis, they are essentially absent in the true vocal cords.

**Hypopharynx**

The pharynx is a hollow muscular tube extending from the skull base to the lower border of the cricoid cartilage. It is arbitrarily divided into three regions: nasopharynx, oropharynx, and hypopharynx. The hypopharynx (also known as laryngopharynx) lies behind the larynx and partially surrounds it on either side, commencing from a plane of the superior border of the hyoid bone (or floor of the vallecula) to the inferior border of the cricoid cartilage. It is continuous with the oropharynx above and with the cervical esophagus below. The junction of hypopharynx with the cervical esophagus corresponds to the sixth cervical vertebra. The lumen of the hypopharynx is cone-shaped, wide superiorly and rapidly narrowing in the postcricoid and cervical esophageal areas. The hypopharynx has three components: right and left pyriform sinuses, postcricoid area, and lateral and posterior pharyngeal walls. The pyriform sinuses are extralaryngeal gutters nestled against the thyroid lamina. Each pyriform sinus is shaped like an inverted pyramid with the apex pointed toward the lower limit of the cricoid cartilage. The superior border corresponds to the pharyngoepiglottic fold. Each sinus has three walls, medial, lateral, and anterior. The postcricoid area forms the anterior wall of the hypopharynx and connects the two pyriform sinuses. It extends from the level of the arytenoid cartilages to the inferior border of the cricoid cartilage (947). The lateral pharyngeal wall merges with the pyriform sinus. The posterior pharyngeal wall extends from the level of the superior surface of the hyoid bone to the inferior border of the cricoid cartilage. The hypopharynx is richly supplied with lymphatics. The major drainage is along the jugular chain (levels II, III and IV), retropharyngeal lymph nodes and the node of Rouvière at the skull base. The hypopharynx is typically lined by nonkeratinizing squamous epithelium, although areas of parakeratin or orthokeratin can be seen secondary to chronic irritation. The lamina propria contains scattered lymphoid aggregates as well as mucoserous glands.

**Trachea**

The trachea extends from the lower border of the cricoid cartilage to the carina and averages 11 cm long in adults, varying roughly in proportion to an individual’s height (1660). It is 20–27 mm transversely and 16–20 mm sagittally (64). There are approximately two tracheal cartilaginous rings per centimeter of trachea, with a total of about 18-22.
Although usually referred to as rings, the cartilages are incomplete posteriorly, and form about two-thirds of a circle. The tracheal cartilages are connected to each other by fibroelastic annular ligaments. Sometimes the first tracheal ring may be fused to the cricoid cartilage. The trachea is continuous with the larynx superiorly and the bronchi inferiorly. Anteriorly, it is intimately associated with the thyroid gland and posteriorly with the esophagus. The trachea is lined entirely by ciliated respiratory epithelium and contains abundant mucous glands in the lamina propria. Posteriorly, the non-cartilaginous or membranous portion of trachea contains smooth muscle. The submucosal lymphatics drain toward the posterior part of the trachea and connect with the paratracheal lymph nodes. They also anastomose with subcarinal, peribronchial and esophageal lymph nodes [64].

**Neck dissections**

A neck dissection is a tissue mass containing the cervical lymphatics. In its classical form, it extends from the submandibular soft tissues to the supraclavicular fatty tissue, laterally bordered by the platysma, and medially by the internal jugular vein. The lymph nodes in this area are divided into 6 different compartments, referred to as levels [2183]. Level I is subdivided in two compartments, the submental area (level IA) that lies between both anterior bellies of the diagastric muscle and the hyoid bone dorsally, and the submandibular area (level IB) that lies between the anterior belly of the diagastric muscle medially and the mandibular bone laterally. Dorsally, this area is bordered by the tendon between the anterior and posterior belly of the diagastric muscle that is attached to the hyoid bone, and the stylohyoid muscle. Thus, the triangle of soft tissue enclosed anteriorly and laterally by the mandible and dorsally by the hyoid is subdivided into one median compartment, the submental area and 2 lateral compartments, the submandibular areas.

Level II represents the upper jugular (cervical) group of lymph nodes. This area extends from the base of the skull superiorly to the level of the inferior border of the hyoid bone inferiorly. The lymph nodes in this area mainly cluster in the vicinity of the internal jugular vein and are laterally covered by the body of the sternocleidomastoid muscle.

Level III represents the middle jugular (cervical) group of lymph nodes. These lymph nodes are located around the middle third of the internal jugular vein that superiorly begins where the upper jugular compartment ends; the lower border lies at the inferior border of the cricoid cartilage.

Level IV comprises the lymph nodes located around the lower third of the internal jugular vein extending from the inferior border of the cricoid cartilage superiorly to the clavicle inferiorly. Level V is the lymph nodes collectively taken together as the posterior triangle group. This is a triangular area lying between the anterior border of the trapezius muscle posteriorly, the posterior border of the sternocleidomastoid muscle anteriorly and the clavicle caudally. It is subdivided into a superior compartment, level VA that contains the spinal accessory lymph nodes and a lower compartment (level VB) that contains the transverse cervical and the supraclavicular lymph nodes. A horizontal plane through the inferior border of the anterior cricoid arch separates both sublevels.

Level VI is the anterior compartment. This compartment has the hyoid bone as its cranial and the suprasternal notch as its caudal border. Both lateral borders are the common carotid arteries. This area is rectangular and lies between the area defined as level I above and the sternum below.

Four different types of neck dissections are recognized [2183].

A radical neck dissection consists of lymph nodes from level I through V. The internal jugular vein, sternocleidomastoid muscle, and spinal accessory nerve also form part of it. A modified radical neck dissection comprises all lymph nodes from levels I-V while preserving one or more of the non-lymphoid structures that should be specified, e.g. modified radical neck dissection with preservation of spinal accessory nerve.

If less than level I-V is removed, the neck dissection is referred to as selective, while specifying the levels that are included. The use of terms such as supra-omohyoid neck dissection is less preferable due to ambiguities about the extent of the surgical procedure. Extended radical neck dissection is the fourth type. This term refers to any type of neck dissection that consists of a radical neck dissection together with additional structures either lymphatic or non-lymphatic that have to be identified specifically. These structures may be additional lymph node compartments, nerves, or blood vessels.

Examination of a neck dissection should be done with the following questions in mind: (1) does the specimen contain lymph nodes with metastatic deposits, (2) if so, how many lymph nodes with metastases are present, specified for each level (3) what is the size of the largest positive lymph node, necessary for staging, and (4) is there extracapsular tumour spread? Dissection of the specimen starts with determination of the type of neck dissection and identification of the various lymph nodes levels and any additional non-lymphoid structures that may have been removed. As the anatomical boundaries that are used by the surgeons to identify the lymph node levels are not present in the specimen, these cannot be used by the pathologist. Optimal processing of a neck dissection therefore requires that the surgeon submit the specimen with all lymph node levels properly labelled.

**Epidemiology**

**Age and sex distribution**

Laryngeal and hypopharyngeal squamous cell carcinoma (SCC) occur most frequently in the sixth and seventh decades, but some cases have been described in children [123,1934]. They are more common in men [344,2113] though the male:female ratio is decreasing in some countries; women are becoming increasingly affected because of increased prevalence of smoking over the last two decades [584]. Tracheal SCC occurs predominantly
between 40 and 60 years of age, men are affected at least twice as often as women [1040].

**Incidence**

SCC comprises about 95% of laryngeal malignancies. The majority originate from the supraglottic and glottic regions, although there are geographic variations in the relative ratio between these two sites. The incidence in men is high (10/100,000 pa or more) in southern and central Europe, southern Brazil, Uruguay and Argentina and among Blacks in the United States. The lowest rates (<1/100,000 pa) are recorded in SouthEast Asia and central Africa. The incidence in women is below 1/100,000 pa in most populations. An estimated 140,000 new cases occurred worldwide in 1990, 86% of these patients were men [1980,1981]. The incidence is slightly more common in urban than in rural areas [344,2113].

There are also geographic differences in the topographic distribution of the laryngeal SCC [126]. In France, Spain, Italy, Finland and the Netherlands, supraglottic SCC predominates, while in the United States, Canada, England and Sweden glottic SCC is more common. In Japan, SCC is approximately equally distributed between the two sites. Interpretation of incidence rates of hypopharyngeal cancer is probably complicated by absence or misclassification within subsites of the pharynx. Recorded incidence is highest among men (>2.5/100,000 pa) in India, Brazil and Central and Western Europe, and is lowest (<0.5/100,000 pa) in East Asia, Africa and Northern Europe. Incidence among women is low (<0.2/100,000 pa) in most populations except India, where rates up to 1/100,000 pa are recorded [1981]. This is probably due to the fact that tobacco is more often chewed than smoked in India.

Tracheal carcinoma is rare with approximately one tracheal carcinoma per 75 laryngeal carcinomas. It accounts for less than 0.1% of cancer deaths [1040]. SCC is the most frequent malignant tumour of the trachea representing 56-73% of all tracheal carcinomas [1040].

**Trends**

The incidence of laryngeal and hypopharyngeal SCC is increasing in much of the world, both in men and in women. This increase is related to changes in tobacco and alcohol consumption [344].

Primary prevention of laryngeal and hypopharyngeal SCC could be achieved by cessation of smoking and reduction of alcohol consumption [344].

**Etiology**

**Tobacco and alcohol - Larynx**

Most cases of laryngeal cancer in Western countries are related to smoking and alcohol abuse [90]. The combined effect follows a multiplicative rather than additive model [285,772,1607,1608,1800,1943,2647,2885]. The increased relative risk (RR) for alcohol consumption differs by site, and is higher for the supraglottis and hypopharynx and lower for the glottis and subglottis [2647]. The impact of increased RR (10x) for smoking is stronger for glottic than supraglottic SCC [2647]. Studies in several populations have shown a direct dose-related response between smoking and SCC and the benefits of cessation. Smoking black tobacco cigarettes entails a stronger risk than smoking blond tobacco [2235]. Other smoking habits that increase the RR of laryngeal SCC include: smoking at a young age, long duration, high number of cigarettes per day, and deep smoke inhalation [195,1035]. The influence of tobacco on RR of laryngeal SCC is confirmed even for non-drinkers [308,2833]. Case controlled studies from Italy and Switzerland show an increased RR of 2.46 for heavy drinkers and laryngeal SCC. The RR for current smokers who do not drink is 9.38 [238]. Avoiding cigarettes and alcohol could prevent about 90% of laryngeal and hypopharyngeal SCC [718].

**Tobacco and alcohol - Hypopharynx**

Studies from India have also reported an association between chewing tobacco-containing products [968] and hypopharyngeal SCC. Tobacco and alcohol are also the main risk factors for hypopharyngeal SCC. The effect of alcohol is stronger and the impact of tobacco is weaker than for laryngeal SCC.

**Asbestos and occupational exposure**

There is controversy regarding occupational asbestos exposure and increased risk for developing laryngeal SCC [247,1255,1982,2484]. A recent review has not supported a causative role for asbestos exposure [283]. However, there is evidence supporting other occupational exposures and increased risk of laryngeal SCC, such as polycyclic aromatic hydrocarbons, metal dust, cement dust, varnish, lacquer, etc [1608]. After adjustment for alcohol and tobacco consumption, the increased risk ranged from 1.8 for cement dust to 2.7 for polycyclic aromatic hydrocarbons. Significant associations are also found with ionizing radiation, diesel exhausts, sulphuric acid mists and mustard gas [1608,2821].
Human papillomavirus (HPV)
There is conflicting evidence implicating HPV16, in 3-85% of laryngeal SCC (1523). The prevailing opinion is that HPV has a minor causative role, if any, in laryngeal carcinogenesis (853,1253, 1510,1523,1999,2330). Additionally, HPV DNA has been detected in 12-25% of individuals with clinically and histologically normal larynges (1912,2172), suggesting that the occasional demonstration of HPV in laryngeal SCC may be incidental.

Diet and nutritional factors
A protective effect is probably exerted by high intake of fruits and vegetables (238, 565,1405,1910,1951,2003,2885,2901). Specific evidence regarding carotenoids and vitamin C, is inadequate for a conclusion (2821). Maté drinking has been suggested to be a risk factor in studies from Brazil and Uruguay (90).

Gastroesophageal reflux
Gastroesophageal reflux has been related to increased risk of laryngeal SCC, especially among patients who lack other major risk factors (80,812,1782, 2724). Gastroesophageal reflux may act as a promoter in the presence of tobacco and alcohol (812).

Genetic susceptibility
There is no evidence of strong genetic factors in laryngeal carcinogenesis; however, polymorphisms for enzymes implicated in the detoxification of alcohol and tobacco, such as alcohol and aldehyde dehydrogenases, are likely to represent weak susceptibility factors, with relative risks in the order of 1.5-2 (200,1590). Bloom syndrome is an inheritable condition with a predisposition towards laryngeal and hypopharyngeal SCC.

Pathology overview and principles
Compartmentally, the supraglottis is distinct from the glottis and subglottis. The supraglottis is embryologically derived from the buccopharyngeal anlage (branchial arches V and VI). The fascial compartmentalization, as well as the lymphatic drainage is distinct for the supraglottis and glottis and is the oncologic basis for the supraglottic horizontal laryngectomy. Dye injected into the supraglottis remains confined and does not travel to the ventricular or glottic tissues. Likewise, glottic dye injections do not pass superiorly to the ventricle or inferiorly to the mucosa overlying the cricoid cartilage. In fact, the mucosa overlying the lamina propria of the glottis (Reinke’s space or laryngeal bursa) may burst from fluid distention rather than allowing injected dye to extend into the ventricle or cross the anterior commissure. These studies also confirm that the larynx is divided into right and left compartments (2087).

The anatomic site of occurrence of tumour within the larynx can influence 1) the type of presenting symptoms, 2) stage at presentation, 3) treatment, and 4) prognosis. The vast majority of malignancies of the supraglottis and glottis are SCC. However the relative distribution of SCC per laryngeal compartment varies worldwide. Non-squamous tumours comprise a small subset of laryngeal malignancies, and are more likely encountered in the supraglottis and infraglottis than the glottis. Glottic tumours present with hoarseness and are typically small when detected. In contrast, the supraglottis is a clinically silent area and, as such, tumours in this site are often large at the time of diagnosis. Epiglottic tumours may present with a change in vocal quality (a muffled or “hot potato voice”), airway obstruction, dysphagia and/or cervical metastasis. Tumours at the base of the epiglottis may escape visualization at indirect laryngoscopy (“Winkelkarzinom” or “cancer in the corner”). Primary ventricular tumours are rare and often remain obscured on laryngeal examination, merely forming a bulge beneath the false vocal cord. Tumours of the pyriform sinus are usually large when discovered and typically present as odynophagia or referred otalgia. If the tumour involves the medial wall or the apex of the pyriform sinus, vocal cord dysfunction may result.

Content of surgical pathology report, including cervical lymph nodes
The surgical pathology report of a laryngectomy specimen should indicate the type of procedure (hemilaryngectomy), and whether any additional tissues are attached (neck dissection, thyroid gland, parathyroid gland). Additional features that should be addressed include 1) site of origin, size and extent of the tumour; 2) histologic type and grade; 3) presence of perineural, lymphovascular, cartilaginous and/or extralaryngeal invasion and 4) status of the resection margins (3,290). The neck dissection should include the details as stated above (see ‘Anatomy - Neck dissections’). Surgical pathology report of a hypopharyngectomy specimen should indicate whether the tumour is arising from the
A detailed history and physical examination are often required. Imaging studies should always precede endoscopy since the latter procedure often results in edema and a decrease in the accuracy of image studies. Direct endoscopy is not only necessary to obtain a biopsy but also important to further evaluate the extent of the disease and rule out additional primary tumours. Distant metastasis frequently distinguishes surgical from non-surgical candidates. Accordingly, tests for hepatic function and imaging studies of lung and bones are invaluable.

### Clinical features and diagnostic procedures

A detailed history and physical examination with attention to specific symptoms will often point to the site and extent of the tumour. There are four critical variables that guide the clinician toward the most appropriate course of therapy: pathologic diagnosis, local extent of tumour, status of regional lymph nodes and presence or absence of distant metastasis.

Imaging studies should always precede endoscopy since the latter procedure often results in edema and a decrease in the accuracy of image studies. Direct endoscopy is not only necessary to obtain a biopsy but also important to further evaluate the extent of the disease and rule out additional primary tumours. Distant metastasis frequently distinguishes surgical from non-surgical candidates. Accordingly, tests for hepatic function and imaging studies of lung and bones are invaluable.

### Second primary tumours

Second primary tumours (SPT) are defined as additional primary malignancies that are distinctly separate from the index tumour (IT). Synchronous SPT are diagnosed at the same time, or within six months of the IT. If the SPT is discovered after six months, it is classified as metachronous. The median prevalence of synchronous SPT for the upper aerodigestive tract (UADT) is 9% (1027). The annual risk for SPT, is rather constant and varies between 1.5% (453) and 5.1% (1309) among patients with UADT SCC. The definition of SPT has been further expanded based on molecular markers of clonality or genetic profiling. This includes comparing patterns of loss of heterozygosity (LOH) and specific p53 mutations at various hotspots among IT, SPT and adjacent mucosa. There are draw backs to comparing only LOH patterns: 1) LOH at various loci can be so frequent, as to be coincidentally present in two tumours, and 2) Progression in genomic loss can be seen with tumour recurrence. This issue is addressed to some extent by p53 mutational analysis, however, identical mutations can occur at various hotspots in 5% of tumours. Some studies have revealed both similarities and discordan ces in genetic profiling between paired IT and SPT (248). Concordant genetic profiles of IT and adjacent mucosa support the concept of mucosal field cancerization as a clonal expansion phenomenon in proximity to the IT. So SPT can arise either as related (clonal) events via lateral mucosal spread of premalignant cells, or as genetically unrelated events. Furthermore, there appears to be an indirect relationship between the distance from the IT to SPT, and the time interval between both, and genetic clonality. Thus synchronous multicentric tumours may be explained by the migration or the distant settling of tumoral cells, whereas distant SPT or metachronous tumours would be better explained by the concept of field cancerization. Newly proposed definitions for “true” SPT, local recurrence, second field tumour and metastasis have been proposed based on molecular profiling (248). “True” SPT are genetically distinct with discordant genetic profiles compared with the IT. If two metachronous carcinomas yield concordant genetic profiles, then the latter tumour is a locoregional recurrence of the former. A “second field tumour” (SFT) distinguishes a second, genetically discordant neoplasm adjacent to the IT, not as a local recurrence but due to the second tumour arising within the same “condemned mucosa” of the IT.

The likelihood and site for developing SPT are influenced by the site of the IT. For UADT IT, the most frequent site for SPT is within the UADT (453,1027,1309,1473,1898); usually an oral SPT associated with intraoral IT. With respect to the larynx, the risk of developing SPT is higher in patients with supraglottic tumours than in those with glottic tumours (1473,1898). Patients with glottic carcinomas are more likely to develop SPT along the respiratory axis (usually lung carcinoma), whereas patients with supraglottic carcinomas are more likely to develop SPT along the aerodigestive axis. No doubt this relates to the specific environmental promoters. The risk of developing SPT clearly correlates with tobacco and alcohol abuse. This risk is more than doubled in patients using tobacco and alcohol, as compared to those patients without exposure (1473). There is a direct dose-dependent relationship between tobacco and alcohol exposure and risk of SPT development (453,1473). The risk of developing a SPT after laryngeal IT increased proportionally to the number of cigarettes smoked per day at the time of the diagnosis (1106). Susceptibility towards the development

<table>
<thead>
<tr>
<th>Histology</th>
<th>N</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carcinoma-in-situ</td>
<td>46</td>
<td>9.6%</td>
</tr>
<tr>
<td>Squamous cell carcinoma</td>
<td>383</td>
<td>79.9%</td>
</tr>
<tr>
<td>Verrucous carcinoma</td>
<td>5</td>
<td>1.0%</td>
</tr>
<tr>
<td>Spindle cell carcinoma</td>
<td>8</td>
<td>1.7%</td>
</tr>
<tr>
<td>Lymphoepithelial carcinoma</td>
<td>1</td>
<td>0.2%</td>
</tr>
<tr>
<td>Subtotal</td>
<td>443</td>
<td>92.5%</td>
</tr>
<tr>
<td>Neuroendocrine carcinoma</td>
<td>4</td>
<td>0.8%</td>
</tr>
<tr>
<td>Adenoid cystic carcinoma</td>
<td>2</td>
<td>0.4%</td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td>3</td>
<td>0.6%</td>
</tr>
<tr>
<td>Chondrosarcoma</td>
<td>8</td>
<td>1.7%</td>
</tr>
<tr>
<td>Liposarcoma</td>
<td>3</td>
<td>0.6%</td>
</tr>
<tr>
<td>Lymphoma</td>
<td>6</td>
<td>1.2%</td>
</tr>
<tr>
<td>Plasmacytoma</td>
<td>1</td>
<td>0.2%</td>
</tr>
<tr>
<td>Malignant granular cell tumour</td>
<td>1</td>
<td>0.2%</td>
</tr>
<tr>
<td>Secondary papillary thyroid carcinoma</td>
<td>7</td>
<td>1.5%</td>
</tr>
<tr>
<td>Fibrosarcoma, low-grade</td>
<td>1</td>
<td>0.2%</td>
</tr>
<tr>
<td>Total</td>
<td>479</td>
<td>100.0%</td>
</tr>
</tbody>
</table>

*Mount Sinai Medical Center, 1994-2003*
of SPT can be demonstrated via mutagen sensitivity tests, such as bleomycin-induced chromatid breaks of cultivated lymphocytes [461,462]. Increased mutagen sensitivity is significantly associated with an increased risk of SPT, with higher risk for both smoking-related and all SPTs (relative risks 2.62 and 2.77, respectively) [2450]. A significantly higher number of chromatid breaks are seen in patients with multiple cancers (mean 1.20) than in patients with a single cancer (mean 0.96) [461,462]. Radiation exposure is also carcinogenic, yet might also have a protective effect on the development of SPT. For patients with laryngeal IT, the latency period for SPT development in irradiated regions was significantly longer than that in non-irradiated patients, suggesting that radiotherapy (RT) may delay the development of SPT [1898]. In patients with laryngeal IT treated by primary RT, the incidence of laryngeal SPT was lower (4.3%) as compared to those patients with laryngeal IT treated primarily surgically, (9.2%), again implying a protective effect [1684].

**Prognosis and predictive factors**

Small glottic or supraglottic SCC can be treated conservatively by laser excision, limited resection, or primary radiotherapy (RT), with curative potential, and overall good survival [1605]. RT failures can be salvaged by conservative, potentially curative voice-sparing surgery. Glottic or supraglottic carcinomas that fix the vocal cord(s) can be treated either by primary resection, with possible adjuvant RT, or organ sparing protocols (neoadjuvant chemotherapy with curative RT). If the carcinoma persists or recurs, overall survival is not compromised by delayed, salvage, total laryngectomy.

The TNM tumour classification consistently correlates, on multivariate survival analyses, with disease-free and overall survival. Among TNM stage IV patients, extensive cartilage invasion and/or bulky tumour volume are predictors of poor response to chemoradiotherapy; these patients are best treated with primary resection and possible adjuvant RT [1883]. Clinical comorbidities have been demonstrated to significantly affect survival over TNM prognosticators [2041]. The Washington University Head and Neck Comorbidity Index incorporates seven conditions (congestive heart disease, cardiac arrhythmia, peripheral vascular disease, pulmonary disease, renal disease, cancer controlled, and cancer uncontrolled) weighted according to severity, and is a significant predictor of survival.

Generally, histological grading has limited impact on survival. By contrast, the pattern of tumour invasion at the advancing host/tumour interface has been demonstrated, by itself or in combination with other histological variables, to have predictive value for laryngeal carcinoma [290]. Thus within T1/T2 laryngeal SCC, biopsy assessment of the pattern of invasion, may be utilized to predict which patients may respond to primary RT, versus which patients are better treated by primary resection. In multivariate analysis, overexpression of p53 is predictive of improved overall survival [1103]. p53 overexpression and elevated PCNA (proliferating cell nuclear antigen) index have been demonstrated to be significant independent predictors of successful organ preservation [249].

**Hypopharynx**

SCC is optimally treated with surgery and adjuvant RT. Prognosis is inversely related to TNM stage and extracapsular spread. [148,1370].

**Trachea**

SCC is the most common primary tracheal malignancy, which is usually treated with RT. Poor prognosticators include mediastinal and distant metastases and poor patient performance status [393,1217].
Squamous cell carcinoma

Definition
Squamous cell carcinoma (SCC) is the most common malignancy of the larynx, pharynx and trachea. It occurs mainly in adult males who abuse tobacco and alcohol, and is characterized by squamous differentiation.

ICD-O code 8070/3

Synonym
Epidermoid carcinoma

Epidemiology
See Introduction page 113-114.

Etiology

Localization
The most common sites for laryngeal SCC vary according to geography, with the supraglottic and glottic regions being the most common locations. Hypopharyngeal SCC originates most frequently in the pyriform sinus, followed by the posterior pharyngeal wall, and the postcricoid area (1053,2661).

Tracheal SCC occurs frequently in the lower third, and less frequently in the upper and middle thirds (1040).

Clinical features
Signs and symptoms
Clinical features depend on the localization of SCC. The most common early symptom in glottic carcinoma is hoarseness. Symptoms of supraglottic, and hypopharyngeal tumours include dysphagia, change in quality of voice, foreign body sensation in the throat, haemoptysis, odynophagia and neck mass. Dyspnoea, and stridor are especially common in subglottic tumours (749). Tracheal SCC usually presents clinically with dyspnoea, wheezing or stridor, acute respiratory failure, cough, haemoptysis, and/or hoarseness (2143).

Macroscopy
Laryngeal and hypopharyngeal SCC may present as a flat plaque with a well-defined, raised edge, or exhibits a polypoid exophytic appearance, which may relate to prognosis. The surface of the tumour is sometimes ulcerated (1711). Tracheal SCC usually presents as a polypoid mass projecting into the lumen, rarely does it grow as a circumferential or an annular mass (1040).

Tumour spread and staging
SCC may spread directly to contiguous structures, or via lymphatic and blood vessels to lymph nodes and more distant sites.

Direct spread to contiguous structures
Supraglottic SCC tends to spread into the pre-epiglottic space, pyriform sinus or towards the base of the tongue, but it rarely invades the glottis and thyroid cartilage. Glottic SCC tends to remain localized for a long period; in late stages of the disease, it may extend to the opposite true vocal cord, to the supraglottis and subglottis; it may also extend through the thyroid cartilage and invade the soft tissue of the neck. The subglottic SCC may spread to the thyroid gland, hypopharynx, cervical esophagus and tracheal wall. SCC that crosses the ventricles and involves the supraglottis and glottis, is termed transglottic SCC (1679). Hypopharyngeal SCC frequently involves the larynx.

Stomal recurrence
Stomal recurrence, defined as recurrent SCC at the mucocutaneous junction of
Squamous cell carcinoma, the tracheostoma, is a well recognized, but infrequent complication after total laryngectomy. Patients with subglottic and postcricoid involvement and advanced stage of the primary SCC are at risk to develop this complication.

**Local and distant metastases**

Laryngeal, hypopharyngeal and tracheal SCC are likely to metastasize to the regional lymph nodes. The location and frequency of lymph node metastases depends upon the site of the primary tumour. Extracapsular spread (ECS) refers to carcinoid penetrating the lymph node capsule and infiltrating extracapsular tissue. Extracapsular spread is further divided into macroscopic and microscopic ECS [337]; macroscopic ECS is evident to the naked eye and appears as matted lymph nodes. Microscopic ECS is only evident on histologic examination and is usually limited to the adjacent perinodal fibro-adipose tissue.

Clinically relevant haematogenous metastases are infrequent but may occur in late stages of the disease. The most common site for spread is the lung, and less commonly, liver and bones [2435]. In patients with blood-borne metastases, regional lymph node metastases are usually also present or have been treated.

**Staging**

Tumours of the larynx and hypopharynx are staged by the TNM system (AJCC and UICC) [947,2418]. Tracheal tumours are not included in the TNM system. In the definition of the N classification, the specified 3.0 cm and 6.0 cm measurements include the total tumour mass in the area (lymph node mass) and not just the individual lymph node size [2801].

**Histopathology**

Squamous differentiation, often seen as keratinization with variable “pearl” formation, and invasive growth are the prerequisite features of SCC. Invasion is manifested by disruption of the basement membrane, and extension into the underlying tissue, often accompanied by stromal reaction. Angiolympathic and perineural invasion are additional signs of malignancy.

The tumours are traditionally graded into well-, moderately-, and poorly differentiated SCC. Well differentiated SCC resembles closely normal squamous epithelium. Moderately differentiated SCC contains distinct nuclear pleomorphism and mitotic activity, including abnormal mitoses; there is usually less keratinization. In poorly differentiated SCC, immature cells predominate, with numerous typical and atypical mitoses, and minimal keratinization. Although keratinization is more likely to be present in well- or moderately-differentiated SCC, it should not be considered an important histological criterion in grading SCC. Most SCC are moderately differentiated, so grading by differentiation is really of limited prognostic value, as compared to pattern of invasion.

**Invasive front**

Tumour growth at the invasive front can show an expansive pattern, an infiltrative pattern, or both. Expansive growth pattern is characterised by scattered small irregular cords or single tumour cells, with poorly defined infiltrating margins and is associated with a more aggressive course [290]. Some guidelines recommend categorizing tumours into cohesive, and non-cohesive fronts, (Reporting Guideline for the Royal College of Pathologists).

**Stromal reaction**

Invasive SCC is almost always accompanied by stromal reaction that consists of desmoplasia with deposition of extracellular matrix and proliferation of myofibroblasts [2907]. Neovascularization is frequently seen.

**Immunoprofile**

SCC expresses epithelial markers such as cytokeratins. The patterns of expression of cytokeratin subtypes may change during malignant transformation and relate to the histologic grade, degree of keratinization, and the likelihood of metastases. Low-grade SCC expresses medium-high molecular weight (MW) cytokeratins, but not low MW cytokeratins [1617]. In contrast, high-grade SCC may express vimentin [2668].

**Electron microscopy**

SCC exhibits desmosomes and attached tonofilaments [880].

**Differential diagnosis**

The diagnosis of SCC is usually not prob-
Tumours of the hypopharynx, larynx and trachea

also includes other malignant tumours, must be distinguished from various sub-
differentiation. Moderately or poorly differentiated SCC lacks prominent branched fibrovascular cores \(171,2602\). Verrucous SCC is composed of papillary projections and bulbous invaginations, lacking cytological atypia. Well-differentiated SCC must also be distinguished from pseudoeplitheliomatous hyperplasia, a benign hyperplastic epithelial condition composed of irregular elongated rete pegs extending deeply into the stroma. It typically occurs in association with chronic infections (tuberculosis, mycosis), trauma, and classically, granular cell tumours. The cytological features of malignancy are not found in pseudoeplitheliomatous hyperplasia.

Distinguishing SCC from radiation changes can be difficult. Radiation can result in ulceration, epithelial and stromal atypia, inflammation and vascular changes. The seromucinous glands may be atrophic. Squamous metaplasia and hyperplasia of ducts, can mimic SCC. Preservation of ductal lumens and lobular architecture aid in making this distinction.

Moderately or poorly differentiated SCC must be distinguished from various subtypes of SCC. The differential diagnosis also includes other malignant tumours, such as adenocarcinoma, neuroendocrine carcinoma, melanoma and lymphoma. The correct diagnosis is best achieved by the use of appropriate immunohistochemistry.

**Precursor lesions**

Precursor lesions are defined as altered epithelium with an increased likelihood for progression to SCC \(847,852,1253,1581\). Epithelial dysplasia is the term used traditionally to describe these microscopic alterations, although other terms have been proposed (see section on epithelial precursor lesions). Pathologists are frequently asked to assess epithelial dysplasia, because it is believed to be an important indicator of malignant potential. The likelihood of malignant change directly relates to the severity of dysplasia. However, it is clear that malignancy can develop from any grade of dysplasia or even from morphologically normal epithelium.

**Histogenesis**

SCC originates from the squamous mucosa or from ciliated respiratory epithelium that has undergone squamous metaplasia.

**Somatic genetics**

**Cytogenetics and comparative genomic hybridization (CGH)**

The most frequent chromosomal alterations detected by CGH are +3q, +5p, +8q, +11q13, +17q, and -3p. Additional alterations, such as +1q, +7p, +7q +9q, +14q, +18p, and -4p, -5q, -11qter, and -18q are also frequent \(1074,1145\). These alterations are very similar to those reported with conventional karyotyping analysis of early passage cells from laryngeal carcinomas \(1219\). Predictive models based on hierarchical branching and distance-based trees indicate +3q21-29 as the most important early chromosomal alteration, followed in importance and chronology by -3p \(1145\). High-level amplifications are found at 3q24-pter and, less frequently 11q13, 18p, 18q11.2, 8q23-24, and 11q14-22. Some of these amplifications are at loci containing known oncogenes \((CCND1 for 11q13)\ \(1074\). Metastasising tumours show a higher number of DNA copy losses than non-metastasising tumours. Losses at 8p, 9q, and 13 are more frequent in metastatic than in primary tumours \(1381\).

**Molecular genetic alterations**

Neoplastic transformation implies modulation of a large number of genes \(1810\) as well as telomerase re-activation as indicated by hTERT expression \(1588\). \(CCND1\) is amplified and overexpressed mostly in advanced cases \(811,1207\). \(MYC\) and \(EGFR\) are amplified in 6-25% of cases although amplification is not related to overexpression \(612,794,811\). Loss of \(RB1\) expression is seen in less than 20% of tumours \(1208,2546\) although LOH at 13q14 is present in 60% or more of tumours, suggesting the existence of other(s) tumour suppressor gene(s) neighbouring \(RB1\) \(2857\).

\(TP53\) mutations are found in 13-50% of laryngeal tumours. The excess of G to T transversions and the codons more frequently affected are both attributed to the carcinogenic effect of tobacco smoking \(1939,2027\). \(TP53\) alterations are found in premalignant lesions indicating participation early in the neoplastic transformation process \(847,1809\). However, neither \(TP53\) overexpression nor \(CDKN1A\) expression are reliable markers for \(TP53\) mutations. Instead, \(CDKN1A\) expression is clearly related to squamous differentiation \(1811\). \(TP73L\), a \(TP53\) homologue with oncogenic potential, maps to 3q27-28, a region with frequent gains. In primary carcioma, low-copy number \(TP73L\) amplification has been detected by fluorescent in-situ
The role of HPV infection in laryngeal carcinoma may be overestimated. The use of PCR-based techniques for the detection of HPV-DNA has yielded variable results. In fact, the virus has even been demonstrated in more than 12-25% of non-neoplastic samples examined. CDKN2A can be inactivated by mutation, homozygous deletion, and promoter hypermethylation. CDKN2A mutations can occur in cases with mRNA and/or protein overexpression. Significant levels of MMP13 mRNA are detectable in some laryngeal tumours, restricted to those that retain features of squamous differentiation. MMP13 expression is coordinated with MMP2 and MMP14 overexpression, two molecules that can efficiently activate MMP13. Both MMP13 expression and MMP14 overexpression are associated with advanced tumours, indicating a more aggressive behaviour. CDH1 expression is lower in metastatic tumours. In addition, gene expression silencing of CDH1 by promoter hypermethylation is more frequent in metastatic (77%) than in primary laryngeal tumours (40%).

Prognosis and predictive factors
The overall 5-year survival rate is 80-85% in glottic SCC, 65-75% in supraglottic SCC, about 40% in subglottic SCC, 62.5% in hypopharyngeal SCC, and 47% in tracheal SCC.

Clinical predictive factors
Stage
TNM remains the most significant predictor of survival.

Localization
Tumour localization is important. The best prognosis has been reported for glottic SCC, and the worst prognosis for hypopharyngeal, subglottic and tracheal SCC.

Other factors that can have an impact on the presentation and outcome of SCC include age, comorbidity (concurrent diseases) and performance status.

Histopathological predictive factors
Resection margins
The complete excision of tumour is the most important principle of oncologic surgery. Negative resection margins are generally associated with decreased recurrence and improved survival. Although controversial, a distance of a few millimeters may be adequate in selected glottic SCC. For supraglottic, advanced glottic, and hypopharyngeal SCC, resection margins have not been precisely defined but distances of at least 5 mm or greater are desired.

Proliferation
Proliferation fraction determined immunohistochemically with antibodies against Ki67 and proliferating cell nuclear antigen (PCNA) have been reported to correlate strongly with the degree of differentiation in SCC and the presence of lymph node metastases. However, proliferation fraction is not an independent prognostic factor.

Lymphovascular and perineural invasion
The penetration of tumour cells into lymphatic and/or blood vessels is associated with an increased propensity for lymph node and/or distant metastases. It tends to occur in aggressive SCC and is associated with recurrence and poor survival. Similarly, perineural invasion is associated with increased local recurrence, regional lymph node metastases and decrease survival.

Extracapsular spread in lymph node metastases
Lymph node metastasis is the single most adverse prognostic factor in head and neck SCC. Recent studies have shown that the presence of extracapsular spread in lymph nodes is strongly associated with both regional recurrence and distant metastases, resulting in decreased survival.

DNA ploidy
The prognostic significance of DNA ploidy has been studied extensively and the results are controversial. Some studies have shown that aneuploid tumours are associated with a higher rate of lymph node metastases and decreased survival while others have not confirmed this. Conflicting results have also been reported regarding the predictive value of DNA ploidy and treatment response.

Genetic predictive criteria
The prognostic value of p53 abnormalities is generally inconclusive for laryngeal carcinoma. CCND1 amplification is related to poor prognosis independent of stage. Simultaneous CDK4 and CCND1 overexpression is associated with poor prognosis. In patients with locally advanced laryngeal cancer, CDKN2A mutations have prognostic significance in predicting adverse outcome.
Verrucous carcinoma

Definition
Verrucous carcinoma (VC) is a non-metastasizing variant of well-differentiated squamous cell carcinoma (SCC) characterized by an exophytic, warty, slowly growing neoplasm with pushing margins.

ICD-O code 8051/3

Synonym
Ackerman tumour [12]

Epidemiology
VC occurs predominantly in men in the 6th and 7th decades of life [1671].

Etiology
VC has been related to tobacco smoking. Human Papillomavirus (HPV) genotypes 16 and 18, and rarely 6 and 11, have been identified in some, but not all, VC [250,289,777,1233,1283].

Localization
Larynx is the second most common site of VC in the head and neck (after oral cavity) and accounts for 15-35% of all VC [1350] and 1-4% of all laryngeal carcinomas [777,1671,1956]. Most arise from the anterior true vocal cords, though it may occur in the supraglottis, subglottis, hypopharynx and trachea [1350,1671].

Clinical features
Hoarseness is the most common presenting symptom; other symptoms include airway obstruction, weight loss, dysphagia, and throat pain [1671,1956]. Enlarged lymph nodes are common and reactive rather than neoplastic [978].

Macroscopy
VC presents as a sharply circumscribed, broad based exophytic warty tumour which is usually firm, and tan to white.

Histopathology
VC consists of thickened club-shaped papillae and blunt intrastromal invaginations of well-differentiated squamous epithelium with marked keratinization and thin fibrovascular cores. The squamous epithelium lacks cytologic criteria of malignancy, and by morphometry, the cells are larger than those seen in SCC [489]. Mitoses are rare, and observed in the basal layers. DNA synthesis (S-phase) is also limited primarily to the basal layers [737]. VC invades the stroma with a pushing, rather than infiltrating border. Dense lymphoplasmacytic host response is common. Intraepithelial microabscesses are seen, and the abundant keratin may evoke a foreign body reaction. The surrounding mucosa shows progressive transition from hyperplasia to VC. A downward dipping of epithelium often "cups" the VC periphery, and is the ideal site for deep biopsy [174,1192]. Hybrid tumours are VC containing foci of conventional SCC. The incidence of hybrid tumours in the larynx is approximately 10% [1956]. It is important to recognize this variant of VC, as it has the potential to metastasize [131,174].

Differential diagnosis
The differential diagnosis of VC includes exophytic SCC, hybrid VC, papillary SCC, keratinizing squamous cell papilloma and verruca vulgaris. VC lacks cytologic atypia, this distinguishes it from exophytic SCC, hybrid VC and papillary SCC. The pushing margins of VC are smooth, in distinction to the irregular shaped invasive islands of SCC. Papillomas have thin, well-formed papillary fronds, with limited keratinization, as compared to the markedly keratinized papillae of VC. Verruca vulgaris of the larynx [722] characteristically contains layers of parakeratotic squamous cells with large keratohyaline granules, identical to their counterpart on the skin.
Prognosis and predictive factors
VC is characterized by a slow, locally invasive growth causing extensive local destruction if left untreated. Pure VC does not metastasize (746,1956). In contrast, hybrid VC has the potential for metastasis and, accordingly, these patients should be managed as similarly staged patients with SCC [1956]. VC has an excellent prognosis; the reported five-year survival rate for laryngeal VC is 85-95% [751,1350]. Patients with VC may be treated by excision (by laser or surgery), or by radiotherapy. Although surgery is more effective, radiotherapy is an acceptable alternative for patients who are poor surgical candidates [978,1350, 1671,1956,2582]. Some reports have suggested that VC may undergo anaplastic transformation following radiotherapy. Critical review of these cases however has shown many to be unrecognized hybrid VC or other carcinomas that were inappropriately labelled as VC.

Fig. 3.14 Larynx verrucous carcinoma. Increased number of cell layers (left), with a broad pushing border of infiltration without cytologically atypical cells (right upper). Keratosis, including parakeratotic crypting is present (right lower).
Basaloid squamous cell carcinoma

Definition
Basaloid squamous cell carcinoma (BSCC) is an aggressive, high-grade, variant of SCC composed of both basaloid and squamous components.

ICD-O code 8083/3

Synonyms
Basaloid carcinoma, adenoid cystic-like carcinoma.

Epidemiology
BSCC occurs in both sexes, but predominantly in men 60-80 years of age {132, 1578, 2128, 2709}.

Localization
The pyriform sinus and supraglottic larynx are the usual sites of involvement {684, 688, 1337, 1578, 1774, 2128, 2709}. It has also been described in the trachea {1992, 1997, 2232}.

Clinical features
BSCC may present with neck mass, hoarseness, pain, sore throat, dysphagia, cough, otalgia, bleeding, and/or weight loss {117, 1997, 2128}.

Etiology
Tobacco and alcohol abuse have been proven to be strong risk factors {117, 132, 1997, 2128}.

Macroscopy
BSCC appears as a centrally ulcerated mass with extensive submucosal induration that may be confused with a minor salivary or soft tissue tumour {132, 1578, 1997, 2128}.

Histopathology
BSCC has two components, i.e. basaloid and squamous cells. Basaloid cells are small, with hyperchromatic nuclei without nucleoli, and scant cytoplasm. They are closely packed, growing in a solid pattern with a lobular configuration, and in some cases, there is prominent peripheral palisading. Comedo-type necrosis is frequent. Distinctive features of BSCC, not found in SCC, are small cystic spaces containing PAS- and Alcian blue-positive material, and stromal hyalinization {117, 2709}. BSCC is always associated with a SCC component which can be either in-situ carcinoma, or invasive keratinizing SCC. The latter is usually located superficially; it may also present as a focal squamous differentiation within the basaloid tumour islands. The junction between the squamous and basaloid cells may be abrupt {117, 132}. Rarely, BSCC is associated with a spindle cell component {1791}. Metastases may demonstrate basaloid carcinoma, squamous carcinoma, or both {688, 1578, 2128}.

Immunoprofile
BSCC expresses cytokeratins and epithelial membrane antigen but the percentage of positive cells is highly variable. To avoid false-negative results, a cocktail of cytokeratin antibodies (i.e. CAM 5.2, AE1/3) is recommended {132}. The antibody 34ßE12, directed against high MW cytokeratins is most sensitive for the detection of basaloid cells {117, 1774}. In the distinction between BSCC and adenoid cystic carcinoma, absence of myoepithelial cells and the presence of dot-like vimentin expression in BSCC can be helpful. S-100 protein reactivity is not helpful in the differential diagnosis, and if observed, usually corresponds to intermingled dendritic cells. BSCC is negative for chromogranin, synaptophysin, and glial-fibrillary acid protein {117, 132, 1337}.

Electron microscopy
Desmosomes and tonofilaments have been observed in basaloid and squamous cells. There are no neurosecretory

Fig. 3.15 Basaloid squamous cell carcinoma of the larynx. A Polypoid tumour with an intact squamous epithelium, subtended by lobules of basaloid cells with areas of central comedonecrosis. B Panoramic view of a tumour that arises from the surface epithelium and is composed of basaloid and squamous nests of cells situated above the epiglottic cartilage.
granules, myofilaments, or secretory granules in BSCC (1082,2709).

**Differential diagnosis**
This includes neuroendocrine carcinoma, adenoid cystic carcinoma, and adenosquamous carcinoma. Neuroendocrine carcinoma typically lacks squamous differentiation, and is strongly positive for neuroendocrine markers. Adenoid cystic carcinoma, especially the solid variant, may resemble BSCC but adenoid cystic carcinoma has a myoepithelial component (132, 1337) and lacks in most instances squamous differentiation. Furthermore, palpable metastatic cervical lymph nodes, quite common in BSCC, are very rare in adenoid cystic carcinoma (2128). Adenocarcinoma and adenosquamous carcinoma can be distinguished from BSCC by the presence of true ductoglandular differentiation, and intracellular mucin.

**Histogenesis**
The suggested precursor of the BSCC is a totipotent primitive cell located in the basal cell layer of the surface epithelium, or in the proximal ducts of minor salivary glands (2128,2709).

**Prognosis and predictive factors**
BSCC is an aggressive, rapidly growing tumour characterised by an advanced stage at the time of diagnosis and a poor prognosis. Metastases to the regional lymph nodes have been reported in two thirds of patients (117,1337,1997,2128), and distant metastases involving lungs, bone, skin and brain, in 35-50% of patients (117,1337,2128). Although controversial, it is generally believed that BSCC is more aggressive than SCC when matched stage for stage (117,736,1337,1578,2709,2787,2798).

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**Fig. 3.16** Basaloid squamous cell carcinoma (BSCC). A Abundant intercellular hyaline globules conferring a cribriform-like pattern. B Nest of basaloid cells with peripheral palisading of the nuclei and a central keratin pearl. C Peripheral palisading of hyperchromatic columnar nuclei and areas of central necrosis. D Hyaline stromal deposits between nests of basaloid and squamous cells. E Central comedo-type of necrosis in a basaloid nest. F Peripherally palisaded nuclei with hyperchromatic nuclei in cells with high nuclear to cytoplasmic ratio. Abrupt keratinization is noted in the centre of this lobule.

**Fig. 3.17** Metastatic basaloid squamous cell carcinoma in a regional lymph node, displaying both basaloid (left) and squamous (right) components.
Papillary squamous cell carcinoma

Definition
Papillary squamous cell carcinoma (PSCC) is a distinct variant of SCC characterized by an exophytic, papillary growth, and a favourable prognosis.

ICD-O code 8052/3

Epidemiology
PSCC occurs predominantly in males in the 6th and 7th decades [501,743,2602].

Etiology
Smoking and alcohol abuse are etiologic factors [501,2602]. Although HPV has been suggested as an etiologic factor, (929} reported prevalence of HPV has varied from 0-48% [171,501,2488]. Its role in the etiology of PSCC is therefore unsettled.

Localization
The larynx and the hypopharynx are among the most common sites of involvement [501,687,743,2488,2602]. Laryngeal PSCC is most often found in the supraglottis, slightly less often in the glottis, and rarely in the subglottis [131, 687,1187,2602].

Clinical features
Hoarseness and airway obstruction are the most common presenting symptoms. Other features include dysphagia, sore throat, cough, and haemoptysis.

Macroscopy
PSCC presents as a soft, friable, polypoid, exophytic, papillary tumour. It frequently arises from a thin stalk, but broad-based lesions have also been described.

Tumour spread and staging
Metastases to the regional lymph nodes may be present, but distant metastases are rare [743]. Lung metastases have been observed in a few patients with laryngeal PSCC [2488].

Histopathology
The tumour is characterized by a predominant papillary growth pattern [2602]. These papillae have thin fibrovascular cores covered by neoplastic, immature basaloid cells or more pleomorphic cells. Commonly, there is minimal keratosis. Foci of necrosis and haemorrhage are frequent. Multiple PSCC or precursor lesions may occur. Stromal invasion consists of a single or multiple nests of tumour cells with dense lymphoplasmacytic inflammation at the tumour-stromal interface. If no stromal invasion is found, the lesion should be called atypical papillary hyperplasia or PSCC in-situ.

Differential diagnosis
This includes squamous papilloma, verrucous carcinoma, and exophytic SCC. Though squamous papilloma and verrucous carcinoma share similar architecture with PSCC, the latter is easily recognized by atypia of the squamous epithelium [743]. Most PSCC strongly express p53 immunohistochemically [2488]. The distinction between exophytic and papillary SCC can be difficult as the criteria for diagnosing exophytic SCC are not clearly delineated [171,2602]. In general, the papillary stalks of PSCC are much better defined than in exophytic SCC.

Precursor lesions
PSCC may evolve from pre-existing papillary mucosal hyperplasia or squamous cell papilloma. Precursor lesions may be solitary or multiple [171,2488].

Prognosis and predictive factors
Patients with PSCC are generally believed to have a better prognosis than those with SCC though reports in the literature are controversial [131,171,1187,2488,2602]. The better prognosis is probably related to limited invasion.
**Definition**
Spindle cell carcinoma (SPCC) is a biphasic tumour composed of a squamous cell carcinoma, either in-situ and/or invasive, and a malignant spindle cell component with a mesenchymal appearance, but of epithelial origin.

**ICD-O code**
8074/3

**Synonyms**
Sarcomatoid carcinoma, carcinosarcoma, collision tumour, pseudosarcoma,

**Epidemiology**
SPCC occurs predominantly in males in the 7th decade of life [1490,2604].

**Etiology**
SPCC has been linked to cigarette smoking and alcohol consumption [1490, 2604] and may develop after radiation exposure [1482,1490,2604].

**Localization**
Larynx is among the most common sites in the head and neck [204]. Less frequently it arises in the hypopharynx [62]. In the larynx, the glottis is most frequently involved [1490].

**Clinical features**
Patients usually present with hoarseness, dysphagia, and/or airway obstruction [1490].

**Macroscopy**
It usually exhibits a polypoid appearance of variable size. The surface is frequently ulcerated. It may rarely appear as an ulcerative infiltrative lesion.

**Tumour spread and staging**
SPCC metastasizes to the regional lymph nodes in up to 25% of cases; but distant dissemination is less common (5-15 %) [1420,1490,2604].

**Histopathology**
The spindle cell component usually forms the bulk of the tumour, which can assume several patterns. Resemblance to fibrosarcoma or malignant fibrous histiocytoma is most common [1490,2604]. Occasional cases can appear less malignant and resemble a reactive fibroblastic proliferation or radiation-induced stromal atypia [62]. Foci of osteosarcomatous, chondrosarcomatous, or rhabdosarcomatous differentiation may be present, particularly in patients with previous radiotherapy (RT) [1420,1490,2604]. Evidence for squamous epithelial derivation can be seen as either in-situ carcinoma or as invasive SCC. Carcinoma-in-situ can be obscured by extensive ulceration. Infiltrating SCC may be focal, requiring multiple sections for demonstration [1482]. Sometimes, only spindle cells are present; in such cases, SPCC can be mistaken for a true sarcoma. (see below). Metastases usually contain SCC alone or both SCC and spindle cell component, and rarely, only the spindle cell component [2457,2604].

**Immunoprofile**
Tumour cells can express both epithelial and mesenchymal markers [1700,2882]. Cytokeratin expression can be demonstrated in spindle cells in 40–85% of cases [671,1700,2407,2535,2610,2882]. The most useful epithelial markers are AE1AE3, CK1, CK18 and epithelial membrane antigen (EMA) [2604]. Spindle cells express vimentin and often other mesenchymal filaments, such as smooth muscle actin, muscle specific actin, and desmin.

**Electron microscopy**
SPCC often displays features of epithelial differentiation in spindle cells, such as desmosomes and tonofilaments [175, 1056,2535,2882].

**Differential diagnosis**
The diagnosis of SPCC generally requires the demonstration of both malignant spindle cells and squamous cell carcinoma, either in-situ or invasive. When a SCC component is inconspicuous, then the spindle cells should be investigated for evidence of epithelial differentiation. However, even in the absence of SCC and negative epithelial markers, SPCC cannot be entirely ruled out. In the larynx and hypopharynx sarcomas are very rare, and SPCC is still more likely. SPCC can also be confused

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**Spindle cell carcinoma**

Figure 3.20 Spindle cell carcinoma. The spindle cell component coexists with basaloid-squamous components seen to arise from the epithelial surface.
with reactive or benign spindle cell proliferations, such as nodular fasciitis, and inflammatory myofibroblastic sarcoma, and low-grade myofibroblastic sarcoma, and myoepithelial carcinoma.

**Histogenesis**
There is mounting molecular evidence that SPCC is a monoclonal epithelial neoplasm (954,2596,2620), with a divergent (mesenchymal) differentiation {172,954, 1490,2596,2620}, rather than a collision tumour, or biphasic derivation, or "pseudosarcoma" {126}.

**Prognosis and predictive factors**
Favourable prognostic features are: low-stage, polypoid rather than endophytic growth, a glottic site of origin, relatively shallow depth of sarcomatoid process, and absence of prior radiation {172}. Although controversial, limited immunoreactivity for polyclonal cytokeratin has been associated with significantly improved survival rates {1944,2604}. The reported 5-year survival is between 65 and 95% {168,2604}.

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![Fig. 3.21 Spindle cell carcinoma. A The ulcerated surface is subtended by a highly pleomorphic spindle cell population with atypical mitotic figures. B Transition between squamous cell carcinoma and spindle cell component. C Pure spindle cell component with marked cellular atypia mimicking a leiomyosarcoma. D Spindle cell component surrounding small nests of squamous cells mimicking synovial sarcoma.](image)

![Fig. 3.22 Spindle cell carcinoma. A The surface epithelium blends imperceptibly with the spindle cell component. B Abrupt areas of squamous differentiation within a spindle cell carcinoma. C Metaplastic cartilage, including malignant transformation (left) and metaplastic osteoid, including malignant transformation (right) can be seen in spindle cell carcinoma. D Strong and diffuse immunoreactivity for keratin is seen in a majority of spindle cell carcinomas.](image)
Acantholytic squamous cell carcinoma

Definition
This is an uncommon histopathologic variant of squamous cell carcinoma (SCC), characterised by acantholysis of the tumour cells, creating pseudolumina and false appearance of glandular differentiation.

ICD-O code 8075/3

Synonyms
Adenoid squamous cell carcinoma, pseudoglandular SCC, SCC with gland-like features, angiosarcoma-like SCC, pseudovascular adenoid squamous cell carcinoma.

Etiology
No special etiological factor has been discovered for the mucosal acantholytic SCC [2876].

Localization
It rarely arises in the supraglottic larynx [1079] and hypopharynx [157], but is more frequent in sun-exposed areas of the head and neck [1847].

Clinical features
There are no special clinical features.

Macroscopy
There are no special gross features.

Tumour spread and staging
As with SCC

Histopathology
This neoplasm is composed of SCC, but with foci of acantholysis in tumour nests, creating the appearance of glandular differentiation. The pseudolumina usually contain acantholytic and dyskeratotic cells, or cellular debris, but they may be empty [157,742]. They are more frequent in the deeper portions of the tumour. There is no evidence of true glandular differentiation or mucin production. The SCC component predominates, and is usually moderately differentiated. Clear and spindle cells may also be present. The stroma is usually desmoplastic, with a lymphoplasmacytic response [157, 742]. The acantholysis may also form anastomosing spaces and channels mimicking angiosarcoma.

Immunoprofile
Acantholytic SCC expresses epithelial markers, such as cytokeratins, and epithelial membrane antigen [742].

Electron microscopy
The tumour cells exhibit hemidesmosomes and attached tonofilaments, and no glandular features thus supporting the squamous origin [2876].

Differential diagnosis
Acantholytic SCC must be differentiated from adenosquamous carcinoma, adenoid cystic carcinoma, and mucoepidermoid carcinoma. This is achieved by demonstrating a lack of true gland formation, absence of myoepithelial cells and negative mucin staining. Vascular markers can be used to distinguish it from angiosarcoma. Cytokeratin, however, might be positive also in some angiosarcomas [939].

Histogenesis
Acantholytic SCC is derived from the surface squamous epithelium.

Prognosis and predictive factors
Prognosis is similar to SCC, however some reports suggest a more aggressive behaviour [157,742,899,2876].

Fig. 3.23 Acantholytic squamous cell carcinoma. A Marked acantholysis of squamous cells giving rise to anastomosing empty spaces with pseudoglandular appearance. B Acantholytic channels, intermingled with dilated capillary blood vessels, mimicking angiosarcoma.
Adenosquamous carcinoma

**Definition**
This rare aggressive neoplasm originates from the surface epithelium and is characterized by both squamous cell carcinoma (SCC) and true adenocarcinoma.

**ICD-O code**
8560/3

**Epidemiology**
There is a male predisposition, with a tendency to develop in the 6th and 7th decades.

**Etiology**
Cigarette smoking and alcohol consumption have been implicated (1294). The role of gastroesophageal reflux has not been well established.

**Localization**
The larynx is the most frequent site, (15,43,831,876,1294); the hypopharynx is occasionally involved (1646,2237).

**Clinical features**
Patients present with hoarseness, sore throat, dysphagia, and/or haemoptysis (1294).

**Macroscopy**
It can present as an exophytic or polypoid mass, or as a poorly defined mucosal induration, frequently with ulceration (876,1294).

**Histopathology**
The main feature is both true adenocarcinoma and SCC. The two components occur in close proximity, but they tend to be distinct and separate, not intermingled as in mucoepidermoid carcinoma. The SCC component can present either as in-situ or as an invasive SCC (43). The adenocarcinomatous component tends to occur in the deeper parts of the tumour. It consists of tubular structures that give rise to “glands within glands”. Mucin production is typically present, either intraluminal or intracellular, and can appear as signet ring cells. However, mucin is not a requirement for the diagnosis in the presence of true glanduloductal formation. Metastases may display both components; one usually predominates.

**Immunoprofile**
There is positive staining for high-molecular weight cytokeratin in both components. The glandular component expresses CEA and low MW cytokeratins: specifically, CK7 is positive, and CK20, is negative (43,1646).

**Electron microscopy**
Features of both squamous and adenocarcinomatous differentiation are found (231,1190).

**DNA ploidy**
A high prevalence of aneuploidy has been demonstrated (43).

**Differential diagnosis**
This includes mucoepidermoid carcinoma.
ma, acantholytic SCC, and SCC invading seromucinous glands, and necrotizing sialometaplasia. The most important differential diagnosis is from mucoepidermoid carcinoma as adenosquamous carcinoma has a poorer prognosis (see Table 3.02) [231]. The presence of mucin in true glandular spaces helps to distinguish adenosquamous carcinoma from acantholytic carcinoma. SCC invading or entrapping mucoserous glands can mimic adenosquamous carcinoma, especially in biopsy specimens. In such cases, preservation of lobular gland architecture, and lack of significant atypia can distinguish SCC from adenosquamous carcinoma. Adenosquamous carcinoma is distinguished from necrotizing sialometaplasia, a benign condition that lacks the cytological features of malignancy. Adenosquamous carcinoma will always have a surface (mucosal) component (dysplasia, in-situ carcinoma), whereas this feature is not seen in mucoepidermoid carcinoma.

**Histogenesis**

Adenosquamous carcinoma originates from basal cells of surface epithelium that are capable of divergent differentiation [1844,1873,2285,2677].

**Prognosis and predictive factors**

Adenosquamous carcinoma is reported as being more aggressive than SCC [876,1844,2237,2611]. Many present as high stage tumours. However, stage for stage comparison with SCC has not been well established. In a recent review, 75% of patients had regional lymph node metastases, and 25% of patients had distant metastases [1294], most commonly to the lungs [731,1294]. The reported 5-year survival rate is 15-25% [831,876,1294]. Half of the patients die of disease after a mean of 23 months (range 12-35 months) [43].
**Definition**
Lymphoepithelial carcinoma (LEC) is an undifferentiated carcinoma with a prominent, reactive lymphoplasmacytic infiltrate, morphological indistinguishable from nasopharyngeal carcinoma.

**ICD-O code** 8082/3

**Synonyms**
Lymphoepithelioma [2317]; undifferentiated carcinoma of the nasopharyngeal type [2317]; lymphoepithelioma-like carcinoma [2741]; undifferentiated carcinoma with lymphoid stroma

**Epidemiology**
LEC of the larynx, hypopharynx and trachea are very rare, and account for less than 0.5% of all cancers in these sites [1722]. There is a male predominance of 4:1, and the mean age is 60 years. In contrast to nasopharyngeal carcinoma, almost all reported cases have occurred in Caucasians [621,1601].

**Etiology**
Smoking and alcohol abuse are noted [621]. Epstein-Barr virus (EBV) is uncommonly demonstrated [621,1601,1637, 2741].

**Localization**
They occur with equal incidence in the larynx and hypopharynx. About two-thirds of the laryngeal tumours are found in the supraglottic region [1601,1637].

**Clinical features**
Patients present with hoarseness, neck mass, sore throat, cough, otalgia, dysphagia or haemoptysis.

**Macroscopy**
The tumour forms a mass that may show deep or superficial ulceration.

**Tumour spread and staging**
Many patients have cervical lymph node metastasis at presentation or early in the course. Distant metastasis (liver, lung, mediastinum, and skin) develops in about one-third of patients [1601,1637].

**Histopathology**
Some LEC show a pure growth pattern, indistinguishable from LEC of other sites. (see Chapter 2 on Tumours of the Nasopharynx). In about half of the cases, there is a component of squamous cell carcinoma that accounts for 10-75% of the entire tumour [1601]. The overlying epithelium can show carcinoma-in-situ.

**Prognosis and predictive factors**
LEC of the larynx and hypopharynx are aggressive, with a propensity for regional lymph node and distant metastasis. A mortality rate of 30% at median follow up of 21 months has been reported [1601].

---

**Fig. 3.27** Lymphoepithelial carcinoma. Islands of undifferentiated carcinoma cells intimately admixed with numerous small lymphocytes and plasma cells.
Giant cell carcinoma

**Definition**
An undifferentiated carcinoma composed of many bizarre multinucleated giant cells, often containing neutrophils or cellular debris in the cytoplasm. It is similar to giant cell carcinoma of the lung (474,768,890,989,2722).

**ICD-O code** 8031/3

**Synonyms**
Large cell carcinoma, pleomorphic carcinoma, undifferentiated carcinoma, anaplastic carcinoma.

**Epidemiology**
Giant cell carcinomas of the larynx are extremely rare (744).

**Etiology**
Smoking and alcoholic consumption have been implicated.

**Localization**
The tumours have all occurred in the larynx, with no site of predilection (744).

**Clinical features**
Progressive dysphonia and dyspnoea are the most common complaints.

**Macroscopy**
The tumours are indistinguishable from squamous cell carcinoma (SCC).

**Histopathology**
The hallmark is the presence of numerous, non-cohesive, bizarre giant cells that contain prominent, frequently multiple nuclei with coarse chromatin and large nucleoli. The cytoplasm is abundant, eosinophilic, sometimes vacuolated, and often contains neutrophils or cellular debris. Additionally, the tumour contains a background population of smaller anaplastic tumour cells. Giant cell carcinoma may exist in a pure or mixed form, in association with SCC, adenocarcinoma, or spindle cell carcinoma (474,768).

**Histogenesis**
The histogenesis remains uncertain, and it has been questioned whether giant cell carcinoma is a specific entity (768).

**Prognosis and predictive factors**
The reported cases have shown a poor prognosis (744).
In a large series of laryngeal tumours from one institution, 72% of salivary gland-type neoplasms were malignant (1039). The majority were mucoepidermoid and adenoid cystic carcinomas.

**ICD-O codes**
- Mucoepidermoid carcinoma: 8430/3
- Adenoid cystic carcinoma: 8200/3

**Mucoepidermoid carcinomas**
Laryngeal mucoepidermoid carcinomas (MEC) are rare, only about 100 cases have been reported (1573). They comprise one third of malignant laryngeal salivary-type tumours (1606). They are much more common in men and most cases present between the ages of 45 and 75 years (peak incidence in the 6th decade) but cases have been reported in children (1750). The most common site is the supraglottis (1937). They present with progressive hoarseness and dysphagia or dysphonia. Nearly half of the cases present with, or develop, cervical lymph node metastases (217,533).

Tumours size is variable. Microscopically they are similar to MEC elsewhere. The behaviour tends to be unpredictable. However, it is acknowledged that a significant number of tumours originally diagnosed as high-grade MEC may have been adenosquamous carcinomas that generally have a poorer prognosis.

**Adenoid cystic carcinoma**
Adenoid cystic carcinoma (ACC) is very uncommon in the larynx and forms only 0.07-0.25% of all laryngeal carcinomas (732,2475), accounting for 1% of all ACC (2446). Only 120 reported cases were identified in a literature review (642). Most occur between the fourth and sixth decades (1039). The majority are subglottic (60%) or supraglottic (35%) and the true cords are involved in only 6% of cases (1573). The microscopic features and outcome of laryngeal ACC are the same as in other sites (469,1942).

**Other salivary-type tumours**
A variety of other carcinomas have been reported in the seromucous glands of the larynx. However, they are very rare and are usually presented as single case reports: acinic cell carcinoma (734,1250, 2146,2454), malignant myoepithelioma (1167), carcinoma ex pleomorphic adenoma (180,232,1729,2220), epithelial myoepithelial carcinoma (1726), salivary duct carcinoma (745,910), papillary adenocarcinoma (2182), mucinous adenocarcinoma (2640) and clear cell carcinoma (1855,2020,2306).

---

**Table 3.3 Salivary gland-type neoplasms of the larynx**

<table>
<thead>
<tr>
<th>Tumour types</th>
<th>No of cases</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pleomorphic adenoma</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>Oncocytic tumours</td>
<td>12</td>
<td>21</td>
</tr>
<tr>
<td>Cystadenoma</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td>11</td>
<td>19</td>
</tr>
<tr>
<td>Adenoid cystic carcinoma</td>
<td>11</td>
<td>19</td>
</tr>
<tr>
<td>Adenosquamous carcinoma</td>
<td>19</td>
<td>33</td>
</tr>
<tr>
<td>Total</td>
<td>57</td>
<td></td>
</tr>
</tbody>
</table>

*Modified from Heffner (1039)
Neuroendocrine neoplasms of the larynx are a heterogeneous group of tumors that vary from benign to highly malignant. Similar to the lung, they can be divided into several types. As a group, they are uncommon with only about 500 cases recorded in the literature as of 1998 (738). The atypical carcinoid is the most frequent, constituting 54% of all neuroendocrine tumors in this site, followed by the small cell carcinoma, neuroendocrine type (34%), paraganglioma (9%) and the typical carcinoid (3%) (125,648,897, 2420,2816).

Cells similar if not identical with Kulchitsky cells of the bronchi are found in the larynx. These, as well as pluripotential endobronchial stem cells, are the putative cells of origin of the typical carcinoid, atypical carcinoid, and small cell carcinoma, neuroendocrine type. Paragangliomas of the larynx are derived from paraganglia normally found in the larynx and are discussed in chapter 8.

**Typical carcinoid**

**Definition**
An epithelial tumor of low-grade malignancy composed of round to spindle cells with histologic, immunohistochemical and ultrastructural evidence of neuroendocrine differentiation.

**ICD-O Code** 8240/3

**Synonyms**
Carcinoid, mature carcinoid, well differentiated (Grade I) neuroendocrine carcinoma.

**Epidemiology**
The typical carcinoid (TC) is the least common of the neuroendocrine neoplasms of the larynx with only 42 cases recorded as of 2005 (2420). It is three times more common in men and most patients are between 45-80 years of age (average 64 years) at diagnosis (738, 2419).

**Table 3.4 Classification of neuroendocrine tumours of the larynx.**

<table>
<thead>
<tr>
<th>Terminology</th>
<th>Synonyms</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. Typical carcinoid</td>
<td>Carcinoid, well differentiated (Grade I) neuroendocrine carcinoma</td>
</tr>
</tbody>
</table>
| B. Atypical carcinoid¹ | Malignant carcinoid, moderately differentiated (Grade II) neuroendocrine carcinoma, large cell neuroendocrine carcinoma 
¹Some atypical carcinomas may fulfill the diagnostic criteria of large cell neuroendocrine carcinoma of lung |
| C. Small cell carcinoma, neuroendocrine type² | Small cell neuroendocrine carcinoma, poorly differentiated (Grade III) neuroendocrine carcinoma 
²Not all small cell carcinomas of the larynx will show neuroendocrine differentiation |
| D. Combined small cell carcinoma, neuroendocrine type, with non-small cell carcinoma, squamous cell carcinoma, adenocarcinoma,etc.) | Combined small cell carcinoma, composite small cell carcinoma |
| E. Paraganglioma | Non-chromaffin paraganglioma |

Fig. 3.30 Typical carcinoid. Typical carcinoid of epiglottis composed of small trabeculae and clusters of cells lying in the lamina propria. The overlying squamous mucosa is intact and free of atypia and/or dysplasia.
Localization
Most occur in the supraglottic larynx in the vicinity of the aryepiglottic fold, arytenoid or false vocal cord.

Clinical features
Symptoms, ranging from three weeks to four years in duration, include dysphagia, hoarseness and a sore throat. At least one patient developed the carcinoid syndrome after the tumour metastasized to the liver [738].

Macroscopy
The tumours have ranged from 0.5 – 3.0 cm. (average 1.6 cm.) in greatest dimension and present as a submucosal or polypoid mass [738].

Histopathology
TCs are composed of round and/or spindle cells that grow in small nests, trabeculae, large sheets, glands and/or rosettes. The cytoplasm is pink and the nuclei have finally stippled or dense chromatin. Nucleoli and mitoses are sparse to absent (less than 2 mitoses/10 HPFs). Necrosis and pleomorphism are not seen. The stroma is highly vascular and often focally fibrotic or hyalinized.

Rarely, carcinoids, either typical or atypical, may be oncocytic or oncocytoid. The distinction depends on the presence (oncocytic) or absence (oncocytoid) of mitochondria on ultrastructural examination. A few may contain mucin and, exceptionally, even amyloid.

Immunohistochemistry
TCs are positive for cytokeratin, epithelial membrane antigen (EMA), carcinoembryonic antigen (CEA), synaptophysin, chromogranin, neuron specific enolase (NSE), and protein gene product 9.5. They are also variably positive for a variety of peptides, including serotonin, calcitonin, bombesin and somatostatin.

Electron microscopy
TCs contain abundant membrane-bound, electron-dense neurosecretory granules varying in size from 90-230 nm [2762]. Cellular junctional complexes are observed as well as numerous mitochondria if the TC is of the oncocytic type.

Differential diagnosis
See under atypical carcinoid.

Prognosis and Predictive factors
Since radiation and chemotherapy are ineffective, surgery is the treatment of choice. The extent of resection should be as conservative as possible, as long as complete removal is achieved. A neck dissection is not warranted. Although the series is small, data indicate that 33% of patients with TCs of the larynx have experienced distant metastases (liver, bones) [648,2419]. At least one patient developed the carcinoid syndrome and another died of disease five years after treatment. While this suggests that TCs of the larynx may be more aggressive than those of the lung, some of these tumours on critical review are probably best classified as atypical carcinoids.
Neuroendocrine neoplasms

Atypical carcinoid

Definition
An epithelial tumour composed of round to spindle cells with histologic, immunohistochemical and ultrastructural evidence of neuroendocrine differentiation exhibiting more mitoses and cellular atypia than a typical carcinoid.

ICD-O code 8249/3

Synonyms
Malignant carcinoid, moderately differentiated (Grade II) neuroendocrine carcinoma, large cell neuroendocrine carcinoma.

Epidemiology
The atypical carcinoid (AC) is 15 times more common than the TC and is the most frequent neuroendocrine neoplasm of the larynx. It is 2-3 times more common in men and has been described in patients from 36-83 years of age (average 61 years) [2769,2816]. Most are heavy smokers.

Localization
Over 90% arise in the supraglottic larynx in the vicinity of the aryepiglottic fold, arynoid or false vocal cord.

Clinical features
Hoarseness, dysphagia, pain in the throat and a neck mass are the usual symptoms. An associated paraneoplastic syndrome is exceptional.

Macroscopy
The tumours present as a tan, gray, pink or haemorrhagic submucosal or polypoid mass 0.2 – 4.0 cm. in greatest dimension (average 1.6 cm) [2769,2816].

Histopathology
ACs are infiltrative tumours that grow in a variety of patterns, including small nests, sheets, trabeculae, glands and/or a combination of these patterns. Cysts with intracystic papillary-like projections of tumour cells may also be seen. In contrast to TCs, the cells are larger and the nuclei are often vesicular and contain prominent nucleoli. Mitoses (usually 2-10/10 HPFs) necrosis, cellular pleomorphism and angiolymphatic invasion are common. Some tumours may even fulfill the diagnostic criteria of large cell neuroendocrine carcinoma of the lung (10 or more mitoses per 10 high power fields and prominent necrosis). Mucinous changes, amyloid, spindle cells and oncocytic-oncocytoid cells may also be observed.

Immunohistochemistry
The tumours may stain for synaptophysin (100%), cytokeratin (96%), chromogranin A (94%), calcitonin (80%), CEA (75%), somatostatin (50%), serotonin (21%), and adrenocorticotropic hormone (17%) [2816].

Electron microscopy
Membrane-bound, electron-dense neurosecretory granules ranging from 70-420 nm are prominent [2769]. Cellular junctional complexes, rough endoplasmic reticulum, mitochondria, Golgi complexes and infrequent bundles of tonofilaments may also be seen.

Differential diagnosis
AC may be confused for a TC, paraganglioma, malignant melanoma, and medullary thyroid carcinoma. The AC is distinguished from the TC by the presence of larger cells, prominence of nucleoli, mitoses, necrosis, pleomorphism and angiolymphatic invasion. AC is positive for cytokeratin, CEA and calcitonin, whereas the paraganglioma is negative for these markers. Malignant melanoma is positive for HMB-45 and tyrosinase and negative for synaptophysin and cytokeratin. Separating AC from metastatic medullary thyroid carcinoma (MTC) may be more problematic since both tumours are positive for synaptophysin, calcitonin and CEA. Clinical and imaging studies to detect the presence or absence of a mass in the larynx or thyroid may offer some assistance. Although the serum calcitonin level is almost invariably elevated in metastatic MTC and usually negative in AC, rare cases of laryngeal AC associated with elevated levels of serum calcitonin have been reported [2409]. Reliance on this test to distinguish between these two tumours is, therefore, not absolute. Knowledge of the serum CEA level (especially if markedly elevated), however, may be helpful. This test is almost universally elevated in MTC, but thus far, has not been reported in association with AC. More recently, thyroid transcription factor – 1 (TTF) has been useful in separating these two tumours. MTC is strongly and diffusely positive for this marker while the AC is typically negative or only focally, weakly positive.

Fig. 3.34 Atypical carcinoid. A Same case as shown in Figure 3.33. Higher magnification shows large cells with prominent nucleoli and mitoses. B Tumour cells are strongly positive for calcitonin.
Factors adversely affecting prognosis include metastatic disease at presentation, positive tumour margins, angiolymphatic invasion, and tumours larger than one centimeter. Determination of DNA ploidy has no prognostic significance.

**Small cell carcinoma, neuroendocrine type**

**Definition**
A highly malignant epithelial tumour composed of small round, oval or spindle cells with evidence of neuroendocrine differentiation.

**Synonyms**
Small cell carcinoma, small cell neuroendocrine carcinoma, poorly differentiated (Grade III) neuroendocrine carcinoma, oat cell carcinoma, anaplastic small cell carcinoma, small cell neuroendocrine carcinoma of intermediate type.

**ICD-O Code**
8041/3

**Epidemiology**
Although the second most common neuroendocrine tumour of the larynx, small cell carcinoma, neuroendocrine type (SCCNET), is still an unusual neoplasm accounting for only 0.5% of all laryngeal carcinomas. It is three times more common in men and is distinctly unusual in patients below 40 years of age.

**Localization**
Although the tumour may arise in any region of the larynx, the supraglottis is, by far, the most common site.

**Clinical features**
Symptoms and signs are those associated with other laryngeal neoplasms and depend on the site of origin. Hoarseness and dysphagia are the usual complaints. Almost half of patients have cervical lymph node metastases at presentation. Exceptionally the tumour may be associated with a paraneoplastic syndrome. Among these include Cushing, Eaton-Lambert, and Schwartz-Bartter syndromes.

**Macroscopy**
The tumours often present as ulcerated submucosal lesions and, as a consequence, may be indistinguishable from ordinary squamous cell carcinoma.

**Histopathology**
The tumour is composed of sheets or rib-
bons of closely packed cells with inconspicuous cytoplasm and round, oval and/or spindle nuclei with dense chromatin and absent nucleoli. Mitoses, necrosis, apoptosis, and lymphatic, vascular, and perineural invasion are common as well as nuclear molding and DNA-coating of the walls of blood vessels. Rare rosettes may be seen. The mucosa is often ulcerated but the marginal epithelium is free of dysplasia. Exceptionally, SCCNET may be associated with a squamous or adenocarcinoma (see “Combined carcinoma” below).

**Immunophenotype**
The immunoprofile is essentially similar to that of the TC and AC. Some may also express thyroid transcription factor – 1.

**Ultrastructure**
Membrane-bound, electron-dense neurosecretory granules ranging from 50-200 nm are scant compared to the TC and AC [2762].

**Differential diagnosis**
The differential diagnosis includes TC, AC, basaloid squamous cell carcinoma, malignant lymphoma, and a metastasis from a primary SCCNET of the lung. Compared to TC and AC, SCCNET is composed primarily of short spindle cells without nucleoli and exhibits more nuclear molding, necrosis and mitotic activity. In addition, SCCNET may be positive for thyroid transcription factor – 1 while the AC is usually negative [1097]. Basaloid squamous cell carcinoma (BSCC) is a biphasic tumour composed of basal and squamous cell components that characteristically grow in a lobular pattern with central comedonecrosis. In addition, BSCCs often exhibit prominent nucleoli, cyst-like areas and hyalinosis and are negative for neuroendocrine markers and TTF. Malignant lymphomas are positive for leukocyte common antigen and negative for neuroendocrine markers. A metastasis from a primary SCCNET of the lung is based primarily on negative imaging studies of the lung.

**Prognosis and predictive factors**
SCCNET is an aggressive tumour with early regional and distant metastasis. Almost half of patients will present with positive cervical lymph nodes and about 60-90% will develop distant metastases, especially to the lungs, liver and bones. The 2- and 5-year survival rates are 16% and 5%, respectively [897]. Because many patients have disseminated disease at the time of diagnosis, radical surgery (laryngectomy with neck dissection) is rarely indicated. Instead a therapeutic protocol using a combination of local radiation and chemotherapy, similar to that for pulmonary SCCNET, is advocated.

**Combined small cell carcinoma, neuroendocrine type**
Small cell carcinoma, neuroendocrine type (SCCNET) associated with a squamous or adenocarcinomatous compo-
Epithelial precursor lesions

Definition
Precursor lesions are defined as altered epithelium with an increased likelihood for progression to squamous cell carcinoma (SCC). The altered epithelium shows a variety of cytological and architectural changes that have traditionally been grouped under the term dysplasia. However, other classifications, e.g. squamous intraepithelial neoplasia (SIN) and squamous intraepithelial lesions (SIL, Ljubljana classification) have also proven to be useful [222,504,846,1054,1055,1253,1254,1711,2317]. Rarely, malignant transformation can develop even from morphologically normal epithelium. Atypia is not considered synonymous with dysplasia. Atypia has been used in the context of inflammatory and regenerative changes particularly referring to cytologic features. In this text, the term atypia refers to cytologic change that may or may not be pre-malignant. Various classifications have evolved to describe the spectrum of histological changes in relation to their malignant potential [222,504,846,1054,1055,1253,1254,1711,2317].

Epidemiology
The entire spectrum of laryngeal and hypopharyngeal precursor lesions are mostly seen in the adult population and affect men more often than women. This gender disparity is especially pronounced after the sixth decade [245]. Mean ages for the first precursor lesion diagnosis are reported from 48.0-56.5 years [243,1253]. The incidence varies worldwide with the magnitude and manner of carcinogen exposure.

Etiology
Precursor lesions are strongly associated with tobacco smoking and alcohol abuse, and especially a combination of these two [221,566,766,1607,1608,1800,2564]. The risk of developing these lesions increases with duration of smoking, the type of tobacco and the practice of deep inhalation. Additional etiological factors are: industrial pollution, specific occupational exposures, nutritional deficiency, and hormonal disturbance [766,1253,1255,1256,1608,1982].

The role of human papillomavirus (HPV) infection in laryngeal carcinogenesis remains unsolved [2412]. The prevalence of HPV in laryngeal carcinoma varies significantly among various studies, ranging from 0% to 54.1% [2517]. Although the overall prevalence of HPV infection found in 9 studies of precursor lesions [97,276,793,853,927,928,1522,2065,2172] was 12.4%, HPV DNA was detected in a clinically and histologically normal larynx in 12-25% of individuals [1912,2172]. Thus, definite evidence of an etiologic role of HPV in precursor lesions, at least at present, is lacking, and HPV infection in precursor lesions may represent an incidental HPV colonization rather than true infection of the laryngeal mucosa.

Localization
Precursor lesions appear mainly along the true vocal cords. Two thirds of vocal cord lesions are bilateral [243]. They can extend over the free edge of the vocal cord to the subglottic surface. An origin in, or extension along the upper surface

Table 3.5 Classification schemas that histologically categorize precursor and related lesions

<table>
<thead>
<tr>
<th>2005 WHO Classification</th>
<th>Squamous Intraepithelial Neoplasia (SIN)</th>
<th>Ljubljana Classification Squamous Intraepithelial Lesions (SIL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Squamous cell hyperplasia</td>
<td>SIN 1</td>
<td>Basal/parabasal cell hyperplasia*</td>
</tr>
<tr>
<td>Mild dysplasia</td>
<td>SIN 2</td>
<td>Atypical hyperplasia**</td>
</tr>
<tr>
<td>Moderate dysplasia</td>
<td>SIN 3***</td>
<td>Atypical hyperplasia**</td>
</tr>
<tr>
<td>Severe dysplasia</td>
<td>SIN 3***</td>
<td>Carcinoma in-situ</td>
</tr>
<tr>
<td>Carcinoma in-situ</td>
<td>SIN 3***</td>
<td>Carcinoma in-situ</td>
</tr>
</tbody>
</table>

* Basal/parabasal cell hyperplasia may histologically resemble mild dysplasia, but the former is conceptually benign lesion and the latter the lower grade of precursor lesions.
** ‘Risky epithelium’. The analogy to moderate and severe dysplasia is approximate.
*** The advocates of SIN combine severe dysplasia and carcinoma in-situ.

Fig. 3.38 Microlaryngoscopic view of laryngeal leukoplakia. Both vocal cords are moderately thickened; an exophytic, well-circumscribed, white plaque is seen in the left vocal cord.

Fig. 3.39 Squamous cell hyperplasia (simple hyperplasia). There is an increased number of ordinary-arranged, otherwise normal cells in the spinous layer. A keratin layer is present on the surface.
of the vocal cord is less common \cite{1253,1332}. The commissures are rarely involved \cite{243}. Hypopharyngeal precursor lesions are rarely identified as the common presentation is established malignancy \cite{2661}. No good data exist regarding tracheal precursor lesions.

**Clinical features**
Most patients with precursor lesions give a history of a few months or more of symptoms, but may be asymptomatic \cite{243}. Symptoms depend on the location and severity of the disease and include fluctuating hoarseness, throat irritation, sore throat, and/or chronic cough. Precursor lesions can be either sharply circumscribed and grow exophytically, or be predominantly flat and diffuse, related in part to the amount of keratin present.

**Macroscopy**
Precursor lesions have a clinically diverse appearance, variously described as leukoplakia (white patch), chronic hyperplastic laryngitis or rarely erythroplasia/erythroplakia (red patch). A circumscribed thickening of the mucosa covered by whitish patches, or an irregularly growing, well-defined warty plaque may be seen. A speckled appearance of lesions can also be present, caused by unequal thickness of the keratin layer. However, the lesions are commonly more diffuse, with a thickened appearance, occupying a large part of one or both vocal cords. Their surface is rough, may be muddy brown to red (erythroplasia), perhaps with increased visible vascularity, or coated with diffuse or dispersed circumscribed whitish plaques (speckled leukoplakia) \cite{1253,1332}. Few white patches are ulcerated (6.5%) or combined with erythroplasia (15%) \cite{243}. Leukoplakia, in contrast to erythroplasia, tends to be well demarcated. In general, leukoplakia has a lower risk of malignant transformation than mixed white and red lesions, or speckled leukoplakia, which has an intermediate risk, and pure erythroplasia which has the highest risk of cancer development \cite{2759}. However, no one clinical appearance is reliably diagnostic of any histologic grade of precursor lesion. Occasionally precursor lesions may appear clinically normal.

**Histopathology**
The epithelium of all precursor lesions is generally thickened. However, in a minority of cases patchy atrophy, thinning of the viable cellular layers, may be present. By definition there is no evidence of invasion. The magnitude of surface keratinization is of no importance. Allocation to categories within each of the classifications requires consideration firstly of architectural features and then of cytology.

1. **Hyperplasia**
Definition: Hyperplasia describes increased cell numbers. This may be in the spinous layer (acanthosis) and/or in the basal/parabasal cell layers (progenitor compartment), termed basal cell hyperplasia. The architecture shows regular stratification and there is no cellular atypia.

**Table 3.6 Criteria used for diagnosing dysplasia**

<table>
<thead>
<tr>
<th>Architecture</th>
<th>Cytology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Irregular epithelial stratification</td>
<td>Abnormal variation in nuclear size</td>
</tr>
<tr>
<td>Loss of polarity of basal cells</td>
<td>Abnormal variation in nuclear shape</td>
</tr>
<tr>
<td>Drop-shaped rete ridges</td>
<td>Abnormal variation in cell size</td>
</tr>
<tr>
<td>Increased number of mitotic figures</td>
<td>Abnormal variation in cell size</td>
</tr>
<tr>
<td>Abnormal superficial mitoses</td>
<td>Increased nuclear-cytoplasmic ratio</td>
</tr>
<tr>
<td>Premature keratinization in single cells</td>
<td>Increased nuclear size</td>
</tr>
<tr>
<td>Keratin pearls within rete pegs</td>
<td>Atypical mitotic figures</td>
</tr>
<tr>
<td></td>
<td>Increased number and size of nucleoli</td>
</tr>
<tr>
<td></td>
<td>Hyperchromasia</td>
</tr>
</tbody>
</table>

**Fig. 3.40** Mild dysplasia (basal-parabasal cell hyperplasia, SIN1). A Note the increased number of basal-parabasal cells with hyperchromatic, uniform nuclei, perpendicularly oriented to the basement membrane. The upper part of the epithelium shows a regular spinous layer and thin parakeratotic layer on the surface. B Increased number of uniform, slightly enlarged basal and parabasal cells, perpendicularly oriented to the basement membrane. Increased number of regular mitoses are evident. At the right corner (lower half) the epithelial cells show minimal cytologic atypia. The upper half of the epithelium is composed of regular spinous cells, which become flattened toward the surface. A thin parakeratotic layer is present on the surface.
2. Dysplasia (intraepithelial neoplasia, atypical epithelial hyperplasia potentially malignant lesions)
Definition: When architectural disturbance is accompanied by cytologic atypia the term dysplasia applies. There is a challenge in the recognition of the earliest manifestations of dysplasia, and no single combination of the above features allows for consistent distinction between hyperplasia and the earliest stages of dysplasia as well as in attempting to rigidly divide the spectrum of dysplasias into mild, moderate and severe categories.

Mild dysplasia
In general architectural disturbance limited to the lower third of the epithelium accompanied by cytological atypia define the minimum criteria of dysplasia.

Moderate dysplasia
Architectural disturbance extending into the middle third of the epithelium is the initial criterion for recognizing this category. However, consideration of the degree of cytologic atypia may require upgrading.

Severe dysplasia
Recognition of severe dysplasia starts with greater than two thirds of the epithelium showing architectural disturbance with associated cytologic atypia. However, as noted in the previous paragraph, architectural disturbance extending into the middle third of the epithelium with sufficient cytologic atypia may be upgraded from moderate to severe dysplasia.

Carcinoma in-situ
The theoretical concept of carcinoma in-situ is that malignant transformation has occurred but invasion is not present. It is not always possible to recognize this morphologically. The following is recommended for the diagnosis of carcinoma in-situ: full thickness or almost full thickness architectural abnormalities in the viable cellular layers accompanied by pronounced cytologic atypia. Atypical mitotic figures and abnormal superficial mitoses are commonly seen in carcinoma in-situ.

Differential diagnosis
Reactive, regenerative or reparative squamous epithelium (for example in response to trauma, inflammation, irradiation or ulceration) may manifest atypical cytology or architectural disturbance. Nutritional deficiencies such as iron, folate, and vitamin B12, can also simulate dysplasia. Such lesions are not considered precursor lesions and should be distinguished from them. Clinical history is helpful, and morphologic changes suggestive of the inciting event (e.g. ulceration, inflammation, haemorrhage, radiation-induced mesenchymal and/or endothelial nuclear enlargement and hyperchromasia) may be present. The epithelial changes in these cases are generally less pronounced than in severe dysplasia/atypical hyperplasia or CIS, atypical mitoses are almost never present, and the epithelium may be thinned, or, if thickened, stratification and maturation often develop as the regenerative/reparative process matures.

Somatic genetics
In studies addressing the genetic changes underlying pre-malignant lesions of the head and neck, the larynx and hypopharynx are often dealt with in a broader anatomic context including the oral cavity. True to current models of carcinogenesis, malignant transformation of the mucosa lining the larynx and other
regions of the head and neck is fundamentally a genetic process that involves activation of key oncogenes and inactivation of critical tumour suppressor genes. These genetic alterations generally occur in order of progression, however, it is fundamentally the net accumulation of multiple genetic alterations that dictates the frequency and pace of progression to invasive carcinoma [318, 319]. Genetic progression does not imply a uniform orderly progression through various stages of histologic progression. By some estimates, progression from normal mucosa to invasive squamous cell carcinoma requires as many as ten independent genetic events [2156]. Loss of heterozygosity studies indicate that the earliest alterations appear to target specific genes located on chromosomes 3p, 9p21, and 17p13 [318]. These alterations, particularly LOH at 9p21, may precede histopathologic evidence of dysplasia [317,2667]. Hyperplasia without any histologic evidence of dysplasia has been found to represent clonal populations of cells sharing the same genetic alterations found in SCC. Alterations that tend to occur in association with higher grades of dysplasia and SCC can include cyclin D1 amplification, pTEN inactivation, and LOH at 13q21, 14q32, 6p, 8, 4q27 and 10q23 [318,787]. Advanced precursor lesions of the head and neck demonstrate a spectrum of genetic alterations that is qualitatively and quantitatively similar to SCC [230, 294,2436].

For some of the chromosomal regions commonly lost or amplified in precursor lesions of the head and neck, the targeted genes have been identified. Two tumour suppressor genes residing at 9p21: p16 (CDKN2/MTS1) inhibit cell cycling via the Rb pathway, and p14(ARF) inhibits cell cycling via the p53 pathway [1022]. The p53 tumour suppressor gene resides at chromosome 17p13. p53 is involved in several cellular regulatory pathways including DNA repair, cell cycle control, and apoptosis [1115]. The cyclin D1 oncogene resides on chromosome 11q13 and is amplified in about a third of SCC [321,811]. However, for most regions of common chromosomal loss such as loss at chromosome 3p, the targeted gene(s) have not yet been well characterized.

Retrospective studies examining the prognostic value of molecular markers, including LOH of chromosomes 3p, 9p21, and 17q13 as well as general aneuploidy, have demonstrated that genetic alterations confer significant risk of malignant progression of precursor lesions. These precursor lesions included clinically defined leukoplakia, with corresponding histologic diagnoses varying along the spectrum of benign to precursor lesions mentioned above. In some cases, retrospective genetic analysis was able to define risk of malignant progression in hyperplastic lesions [1627,2201,2492].

**Prognosis and predictive factors**

Some precursor lesions are self-limiting and reversible, others persist and some progress to SCC [503]. The histopathologic degree of severity of these lesions can be a predictive factor [222,846,1054,1689]. Simple and basal/parabasal cell hyperplasias have a minimal likelihood of malignant progression (0.9%). These patients do not require close clinical follow-up. Lesions classified as atypical hyperplasia (moderate to severe dysplasia) have a 11% rate of malignant transformation [1054]. Diagnosis of precursor lesions implies a need for close follow-up and complete excision depending on the clinical situation [846,1054]. Patients with carcinoma in-situ require more extensive management, depending on the clinical circumstance [504,1253, 1808,2151,2432].
Papilloma / papillomatosis

**Definition**
Squamous cell papillomas are the most common benign epithelial tumours of the larynx, caused by HPV infection. The tumours can be multiple and often recur.

**ICD-O codes**
Papilloma 8050/0
Papillomatosis 8060/0

**Synonyms**
Recurrent respiratory papillomatosis (RRP), laryngeal papillomatosis, juvenile papillomatosis, adult papillomatosis.

**Epidemiology**
They rarely appear as solitary lesions, more frequently, especially in children, as recurrent respiratory papillomatosis (RRP). It is characterized by multiple contiguous lesions with great propensity for local recurrences. There is a bimodal age of distribution; the first peak is before the age of 5 (juvenile form), with no gender predominance. The second peak occurs between the ages of 20-40 years (adult form) with a 3:2 male predominance {178,586,1282,1519}. Its true incidence and prevalence are uncertain. A wide range in incidence, 0.4-4.3 /100,000, has been recorded for various regions worldwide {73,585,1253,1520}.

**Etiology**
HPV-6 and 11 are the most frequent genotypes associated with RRP as well as solitary papillomas {10,848,1484,1788,2074,2101,2411}. Rarely, HPV 16, 18, 31, 33, 35 and 39 have been identified in RRP {389,2004,2074,2173,2461}. The mode of HPV infection in children is perinatal vertical transmission related to maternal genital infection {178,586,2311}. An epidemiological triad has been identified: the first-born child, vaginal delivery and teen-aged mother correlating with juvenile RRP {1282}. Caesarean delivery has not been found to be entirely protective against the disease {2357}. Factors that influence the conversion of HPV exposure to active HPV infection resulting in epithelial proliferation are not known {944}. The mode of HPV infection in adults remains unclear. The reactivation of a latent infection acquired perinatally or adult acquired infection with orogenital contacts have been suggested {10,1281}.

**Localization**
The disease almost invariably involves the larynx, especially true and false vocal cords, subglottic areas and ventricles {10,126}. Papillomas may spread to other laryngeal sites; the most frequent sites of extralaryngeal spread are the oral cavity, followed by trachea and bronchi. Extralaryngeal extension of RRP has been identified in 30% of children and in 16% of adult patients {240}. Endobronchial and pulmonary dissemination occurs in 5% of patients with RRP {585}. Rare cases of isolated tracheal lesions without laryngeal involvement have been reported {2662}. The distribution of RRP follows a predictable pattern, occurring mainly at anatomic sites in which ciliated and squamous epithelia are juxtaposed. An injury of ciliated mucosa after surgical procedures may result in squamous metaplasia creating an iatrogenic squamous-ciliary junction, thereby inducing a new background for additional tumours {1280}.

**Clinical features**
Squamous papillomas have been traditionally divided into juvenile and adult groups {586,1282,1519,2579} and additionally into multiple or solitary groups {1519,1524}. Although caused by the same viruses, they follow distinctly diverse and variable clinical courses. For children, extensive growth with rapid recurrences is characteristic. In some patients, RRP becomes indolent as the patient reaches adulthood. Disease progression is more frequent in the juvenile form, usually associated with subglottic papillomas and prior tracheotomy {2743}. Tracheobronchial extension of RRP is associated with morbidities such as pneumatoceles, lung abscesses, tracheal stenosis and rarely malignant transformation {219}. The relatively small airway in children predisposes to airway obstruction. The clinical course in adults is usually not so dramatic, the development of respiratory distress and other complications are rare, although RRP can be aggressive with multiple recurrences {240,2074}.

**Signs and symptoms**
Adult laryngeal papillomas cause hoarseness {240,1253}. The disease is usually self-limiting. Most children with laryngeal RRP present with dysphonia and stridor, less commonly with chronic cough, recurrent pneumonia, dyspnœa, and acute lifethreatening events {178,240,587}. Delay, due to mistaken diagnoses, such as bronchitis, asthma, and other allergic manifestations, may lead to gradual respiratory distress and urgent tracheotomy, which is associated with more frequent extension into the tracheobronchial tree. The overall mortality rate of patients with

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**Fig. 3.43** Laryngeal papillomatosis. **A** Recurrent respiratory papillomatosis fills the endolaryngeal space. **B** Multilobulated grape like clusters of papillomas are located on the right side and anterior commissure of the larynx. (Endoscopic view)
RRP ranges from 4-14% [126], often due to asphyxia, infection, pulmonary complications, and malignant transformation [109,126,2437].

**Macroscopy**
They are exophytic, branching, pedunculated or sessile masses, pink or red, with finely lobulated surface, disposed to bleeding with minor trauma, presenting either singly or in clusters.

**Histopathology**
Squamous papillomas are composed of finger-like or frond-like projections of squamous mucosa, containing thin fibrovascular cores. Secondary or tertiary branching of papillae may be present. Keratosis is minimal. Frequently, parabasal cell hyperplasia, usually extending up to the mid-portion with a perpendicular orientation of the cells to the basement membrane, is seen. Mitotic features may be prominent within this area. Koilocytosis is occasionally evident [2066]. Premature and abnormal keratinizing individual cells (dyskeratosis) if present, contributes to a disorganized appearance. Premalignant changes are not commonly observed, and should be reported when present [1253].

**Histogenesis**
RRP mainly originates from the squamous epithelium where ciliated and squamous epithelia are juxtaposed [1280]. HPV enters the basal cells through a microtraumatized squamous epithelium. Viral replication occurs in the spinous layer, causing a disturbance of epithelial maturation [2917].

**Prognosis and predictive factors**

**Clinical criteria**
The clinical course of RRP is unpredictable, characterised by periods of active disease and remissions. The presence of HPV in apparently normal mucosa is thought to be a virus reservoir and the source of repeated recurrences [2171,2411]. RRP in the neonatal period is associated with poor prognosis, with a greater need for tracheotomy and likelihood of mortality [587].

**Histopathological criteria**
Significant histological prognosticators of local recurrences and malignant transformation have not been identified [501, 908, 2100].

**DNA ploidy, proliferation**
DNA aneuploidy and Ki-67 proliferative index, in contrast to histologic indices, have been found to predict disease recurrence and extension for children with RRP [2470,2471].

**HPV genotype**
HPV 11 and 16 are associated with more aggressive disease, with frequent recurrence and progression in children and adults [2074,2110,2174].

**Malignant transformation**
Malignant change is not common, but occurs in the setting of smoking, irradiation, or other promoters [2112]. It is an exceptional event in the absence of predisposing factors [196,959,2144,2356]. The overall incidence of cancer development for irradiated patients is 14% and 2% for the non-irradiated [126]. Malignancies occur preferentially in the tracheobronchial tree in children, and in the larynx in adults [944]. Prognosis in children is poor [2422]. HPV 11 is most frequently associated with malignant transformation of RRP [487,1465,1521, 2112], followed by HPV 16 [619] and HPV 18 [2226].

**Fig. 3.44** Squamous papillomas of the larynx. A Higher magnification of a papillary branch. B Pronounced koilocytosis is evident in the upper part of the squamous epithelium. C Positive in-situ hybridization signal for HPV genotypes 6 and 11 in the upper part of the squamous epithelium.

**Fig. 3.45** Branch of a laryngeal squamous papilloma. A Note atypical hyperplasia of the covering epithelium. B Note basal and parabasal cell hyperplasia of the covering epithelium extending up to the half of the epithelial thickness.
In the larynx, benign salivary gland-type tumours are rare and less frequent than the malignant varieties (1039).

**ICD-O codes**
- Pleomorphic adenoma 8940/0
- Oncocytic papillary cystadenoma 8290/0

**Pleomorphic adenoma**
Most pleomorphic adenomas arise in the epiglottis or aryepiglottic folds and can reach several centimetres before producing symptoms. Microscopically, laryngeal pleomorphic adenomas are similar to those in other minor salivary glands.

**Oncocytic papillary cystadenoma (OPC)**
Synonyms: oncocytic cyst, oncocytic papillary cystadenomatosis of the larynx, oncocytic adenomatous hyperplasia, oxyphil adenoma, oncocytoma and adenolymphoma in laryngocele (1641,2845).

Laryngeal oncocytic lesions usually consist of unilocular or multilocular cysts lined by cytologically bland oncocytic epithelium with or without intraluminal papillary ingrowths (748,1548). These lesions probably represent duct hyperplasia and metaplasia rather than true neoplasia. Most patients are older than 50 years, and present with hoarseness or other symptoms. The most frequent locations are the false vocal cords and the laryngeal ventricular areas (850). The lesions are not encapsulated, may be multicentric, and can have a Warthin-like lymphoid component (792). Solid oncocytomas of the larynx resembling those seen in major salivary glands are rare to absent. Recurrence is uncommon and they have no malignant potential.
Malignant soft tissue tumours

**Fibrosarcoma**

**ICD-O code** 8810/3

In the past, the term “fibrosarcoma” was often applied indiscriminately to any malignant spindle cell tumour associated with collagen production. On critical review of these cases, supplemented with immunohistochemistry, it has become apparent that many alleged fibrosarcomas are examples of other entities [1636,2191]. With the possible exception of radiation-induced tumours, de novo fibrosarcoma is now recognized as a relatively uncommon tumour [1831]. The main differential diagnoses include spindle cell carcinoma and, occasionally, inflammatory myofibroblastic tumour, posttraumatic spindle cell nodule, and radiation-induced stromal atypia [2604, 2889].

**Malignant fibrous histiocytoma (MFH)**

**Definition**

An aggressive, highly controversial malignant mesenchymal neoplasm composed of primitive round to spindle-shaped cells, often with admixed inflammatory and multinucleated giant cells, that grows either focally or diffusely in a storiform pattern.

**ICD-O code** 8830/3

**Epidemiology**

MFH of the larynx is an uncommon tumour. It occurs in all age groups (6-68 years) and is more common in males by a ratio of 3:1 [1985,2283].

**Etiology**

Other than those related to prior radiation exposure, there are no known predisposing factors.

**Localization**

MFH is distinctly unusual in the hypopharynx and trachea. In the larynx, the glottis is the site of predilection.

**Clinical features**

Symptoms vary according to location and include hoarseness, airway compromise, dysphagia or a sensation of a foreign body in the throat.

**Macroscopy**

The tumours are sessile to polypoid, firm, often ulcerated and have a yellow-tan to grey-white cut surface.

**Histopathology**

The histomorphology is highly variable but includes several of the following features: histiocyte-like cells, spindle-shaped cells, foam cells, pleomorphic multinucleated giant cells, typical and atypical mitoses, and necrosis. The tumour characteristically grows in a storiform pattern, either focally or diffusely.

**Differential diagnosis**

MFH must be distinguished from spindle cell carcinoma, which may be difficult on small biopsies. In spindle cell carcinoma, the tumour cells are typically positive for cytokeratin, as opposed to MFH. The presence of dysplasia or carcinoma in situ in the overlying mucosa also indicates a carcinoma.

**Prognosis and predictive factors**

Surgery is the treatment of choice. The role of irradiation and chemotherapy is largely untested. In the absence of enlarged lymph nodes, a prophylactic neck dissection is not indicated [2283]. The tumour is unpredictable but certainly has the potential for local recurrence, haematogenous metastasis and death from disease.

**Liposarcoma**

**ICD-O code** 8850/3

Primary liposarcomas of the larynx are rare, comprising less than 20% of all head and neck liposarcomas and fewer than 0.5% of all laryngeal neoplasms. Patients of all ages are affected, with a median of 64 years. There is a marked male to female predominance (nearly 10:1). The tumours, which occur almost exclusively in the supraglottic larynx or hypopharynx (pyriform sinus), most commonly cause airway obstruction. Imaging, especially with MR or CT, will document the lipomatous nature and extent of the mass. The tumours are firm, polypoid pedunculated, up to 10 cm in greatest dimension and demonstrate a lobulated, glistening, translucent cut surface often traversed by bands of fibrous tissue. The mucosa is usually intact. The majority of cases are well-differentiated lipoma-like liposarcomas (grade I), similar to their histologic counterparts in other anatomic sites, with infrequent reports of myxoid and pleomorphic types. Lipoblasts may be scanty necessitating multiple sections. Atypical cells, scattered lipoblasts, and infiltrative growth pattern differentiate liposarcomas from lipomas. In spite of surgical treatment, multiple recurrences are not uncommon (80% of patients). Metastasis has not been reported and the long-term prognosis is excellent (90% 5-year survival) [733,918,1155, 1371,2765,2772].
Leiomyosarcoma

ICD-O-code 8890/3

Leiomyosarcomas arising in the larynx are exceedingly rare accounting for less than 0.1% of all laryngeal malignancies. They present mainly in adults with no gender predilection. Symptoms are non-specific. Tumours can occur anywhere in the larynx but supraglottic lesions have been more frequently reported. They have a histology similar to leiomyosarcomas in soft tissues and demonstrate increased cellularity, nuclear and cellular pleomorphism, cytoplasmic vacuolization, necrosis, hemorrhage, and increased mitotic activity in addition to invasive growth. The diagnosis of leiomyosarcoma requires histologic, immunophenotypic (desmin, actins), and/or ultrastructural (parallel actin filaments, dense bodies and pinocytotic vesicles) confirmation as spindle cell carcinoma must always be excluded. Primary treatment is surgical. A variable prognosis is achieved [840,1247,1530,1635,1668,1969,2208,2373,2706].

Rhabdomyosarcoma

ICD-O code 8900/3

Rhabdomyosarcomas of the hypopharynx, larynx and trachea are poorly documented and exceedingly rare, comprising no more than 2% of all rhabdomyosarcomas [25,518,1084,1293,1508,2215,2781]. They have been described in all age groups and, in the larynx, are centered around the glottic region. Possibly because of early presentation, prognosis has generally been good but death from disease has been recorded. The differential diagnosis includes a spindle cell carcinoma [915].

Angiosarcoma

ICD-O-code 9120/3

Primary angiosarcomas of the larynx are exceedingly rare, with only a few well-documented reports. Despite the fact that nearly 50% of all angiosarcomas occur in the skin and superficial soft tissues of the head and neck, angiosarcoma accounts for less than 0.1% of all head and neck malignancies. Laryngeal angiosarcoma is twice as frequent in men with a mean age at presentation in the 7th decade of life. Symptoms are non-specific; previous radiation exposure is frequently noted. The supraglottis is affected more frequently, specifically the epiglottis, where an increasing size is associated with a worse clinical outcome. Tumours demonstrate the typical histomorphologic features of angiosarcoma in other soft tissue sites. Tumour cells are consistently positive with Factor VIII-related antigen, CD34, and CD31. Contact ulcer, haemangioma, acantholytic squamous cell carcinoma and mucosal malignant melanoma are the principle differential diagnostic considerations. Surgical excision is the treatment.

Fig. 3.49 Rhabdomyosarcoma. A Cells with eosinophilic and striated cytoplasm contain eccentrically placed nuclei with prominent eosinophilic nucleoli. Degeneration is noted. B The neoplastic cells are strongly immunoreactive with desmin, accentuated in both the 'epithelioid' and spindled cells.

Fig. 3.50 Laryngeal angiosarcoma. A Intermediate power demonstrating the anastomosing vascular channels lined by atypical endothelial cells. B A high power showing nuclear atypia of the hobnailed endothelial cells in a laryngeal angiosarcoma. C Factor VIII R-Ag reacts strongly and diffusely with the neoplastic endothelial cells.
of choice, with radiation therapy as necessary, yielding an overall patient survival of about 50% at 2 years. Large tumour size and previous radiation portends a worse prognosis [1556,1692,2044,2055,2280,2591,2847].

**Kaposi sarcoma**

ICD-O-code 9140/3

Kaposi sarcoma (KS) of the larynx is uncommon and only a few well-documented cases have been reported since 1983, coincident with the time frame during which HIV and AIDS were beginning to be recognized. This finding lends support to the strong association of KS of the larynx with the advanced HIV disease in epidemic AIDS rather than an association with the iatrogenic immunocompromised transplant, the endemic African, or the sporadic form. Men are almost exclusively affected, usually in the middle decades of life, presenting with upper airway obstruction. A flat to raised, violaceous, plaque-like mass is usually identified in the supraglottis, although glottic lesions are also frequent. Multifocal involvement is reported. The cut surface is fleshy and demonstrates recent and old haemorrhage. Biopsy is contraindicated, as brisk haemorrhage will require emergent tracheostomy and possible death by exsanguination. Treatment is generally nonsurgical, encompassing radiotherapy or chemotherapy (systemic or intralesional). Laryngeal KS is usually non-lethal [191,499,815,1487,1753,2262,2552].

**Peripheral nerve sheath tumours (PNST)**

ICD-O-codes

<table>
<thead>
<tr>
<th>Schwannoma</th>
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<tr>
<td>Neurofibroma</td>
<td>9540/0</td>
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<tr>
<td>Malignant peripheral nerve sheath tumour (MPNST)</td>
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Both benign (especially schwannoma and neurofibroma) and malignant peripheral nerve tumours (MPNST) can occur in the larynx, although vanishingly rare. Any age group can be affected and there is no gender predilection. Association of neurofibroma with neurofibromatosis-1 (NF1) has been reported in the larynx [385,2508].

Macroscopically, benign and malignant PNST most often involve the supraglottis in relation to the superior laryngeal nerve. The tumours are of variable size and present as smooth, round or lobulated to fusiform submucosal masses, often demonstrating cystic or mucinous degeneration. The mucosa is usually intact, although larger tumours may ulcerate. Schwannoma is encapsulated and solitary with the nerve of origin attached peripherally. In contrast, neurofibroma is non-encapsulated, occasionally multiple or plexiform, especially when it is associated with NF1 and expands the nerve in a fusiform fashion rather than pushing it aside. MPNST are infiltrative, mitotically active and often ulcerate the mucosa. The tumour cells are immunoreactive, often patchy, with S-100 protein, vimentin, epithelial membrane antigen (EMA) and Leu-7. Distinguishing benign from malignant tumours on small biopsies may be difficult [385,659,1202,1235,1974,2508,2591].

**Synovial sarcoma**

ICD-O-code 9040/3

Primary synovial sarcoma of the larynx and hypopharynx is rare, while secondary involvement by direct extension from the neck is slightly more common. Although all age groups may be affected, most patients are young and there is no gender bias. Symptoms are non-specific. The tumours are often exophytic or pedunculated, and infiltrative with surface ulceration. Both monophasic and biphasic synovial sarcomas have been described and are similar to the counterpart in the soft tissue. Immunohistochemically, both epithelial and spindle cells may be reactive with cytokeratin and epithelial membrane antigen (EMA), while only the spindle cells are positive for vimentin. Molecular studies reveal a characteristic translocation t(X;18)(p11.2;q11.2). The prognosis is variable but tends to be better than those arising in soft tissue [573,740,1735,1780,2102].
Inflammatory myofibroblastic tumour

Definition
Inflammatory myofibroblastic tumour (IMT) is a distinct borderline lesion composed of myofibroblastic cells with a variable admixture of inflammatory cells, including mature lymphocytes, histiocytes, plasma cells and eosinophils, and collagen. It occurs predominantly in the soft tissue and viscera but also in the head and neck [2761].

ICD-O code 8825/1

Synonyms
Inflammatory pseudotumour, plasma cell granuloma, plasma cell pseudotumour, pseudosarcomatous lesions/tumours.

Epidemiology
Head and neck IMTs are rare. There are very few comprehensive studies detailing their clinicopathologic features. In contrast to soft tissue and visceral IMTs, that occur predominantly in children and young adults, IMTs of the upper aerodigestive tract are more common in adults (median age of 59) and men [2761].

Etiology
The etiology of IMT is unknown. Previous trauma, and immunosuppression have been implicated [858,2761]. Recently, human herpesvirus-8 DNA sequences and overexpression of interleukin 6 and cyclin D1 have been reported in IMTs [920].

Localization
In the head and neck, inflammatory myofibroblastic tumours are most common in the larynx [396,686,1648, 2761] especially in the region of true vocal cord [2761]. Non-laryngeal sites include the oral cavity [632,677,1531], tonsil [858,2736] parapharyngeal space [383,1165,2739], sinonasal tract [1278, 1616,2165,2429,2739], salivary glands [1178,2739,2794] and trachea [53].

Clinical features
Laryngeal IMTs present with hoarseness, stridor, dysphonia, and/or a foreign body sensation in the throat [2761]. Constitutional and/or systemic signs and symptoms such as fever, weight loss, pain, malaise, anaemia, thrombocytosis, polyclonal hyperglobulinemia and elevated erythrocyte sedimentation rate seen in association with soft tissue and visceral IMTs are not usually a component of upper aerodigestive tract IMTs; however, they are occasionally reported [383, 2165].

Macroscopy
The gross appearance of laryngeal IMT is a smooth, polypoid or nodular lesion with fleshy to firm consistency and varying dimensions.

Histopathology
IMTs are characterized by a submucosal, loose, cellular proliferation of spindled to stellate cells arranged in a storiform to fascicular pattern with a variable component of inflammatory cells. The surface epithelium may be intact, hyperplastic or ulcerated and show reactive epithelial atypia. The inflammatory infiltrate is comprised of mature lymphocytes, histiocytes, plasma cells, eosinophils and scattered neutrophils. The stroma is highly vascularized and ranges from oedematous to fibromyxoid to collagenous. The vascular component varies from widely dilated medium sized vascular channels to narrow, slit-like blood vessels that can be obscured by the myofibroblasts and inflammatory cells. Vascular thrombosis is not present. This overall appearance is similar to a reactive process resembling granulation tissue or nodular fasciitis. The myofibroblasts are spindled to stellate with enlarged round to oblong nuclei, variable nucleoli and abundant, eosinophilic to basophilic appearing fibrillar cytoplasm. In some examples, the myofibroblasts have a more epithelioid or histiocytoid appearance, including round to oval nuclei, prominent nucleoli and abundant, eosinophilic to basophilic appearing fibrillar cytoplasm. The myofibroblasts may also appear as slender axonal (spider-like) cells with elongated nuclei, inapparent nucleoli and long cytoplasmic extensions.
creating a tadpole-like appearance. The myofibroblasts invariably maintain a low nuclear-to-cytoplasmic ratio. Focal nuclear pleomorphism may be present. Mitoses are common, sometimes even numerous, but never atypical. Necrosis and marked nuclear pleomorphism are not seen.

**Immunoprofile**
IMTs show strong diffuse cytoplasmic immunoreactivity for vimentin, and usually variable expression of smooth muscle actin and/or desmin. Cytokeratin staining may be seen but usually focal to absent.

**Electron microscopy**
The tumour cells show myofibroblastic and fibroblastic differentiation [707, 2761].

**Genetics**
Recent evidence reveals the presence of anaplastic lymphoma kinase (ALK) gene rearrangements and expression in IMTs [486, 949, 1440, 2865]. These rearrangements are common in IMTs of children and young adults [949, 1440, 2486] and are uncommon over the age of 40 [367, 486, 1440]. Both the gene rearrangements and protein activation are restricted to the myofibroblastic component, while the inflammatory cell component is normal [270, 466, 466, 486, 949, 1440]. Fusion of ALK to Ran-binding protein 2 gene in IMTs expand the spectrum of ALK abnormalities seen in IMT further confirming the clonal, neoplastic nature of IMTs [1593].

**Prognosis and predictive factors**
Laryngeal IMT is usually cured by conservative resection [2761]. Corticosteroid and nonsteroidal anti-inflammatory agents have been used for treatment resulting in regression in some patients [795, 2487]. A recurrence rate of approximately 25% has been reported for extrapulmonary IMTs [467]. IMT was originally believed to be a reactive non-neoplastic lesion. This has been refuted by the above genetic studies. An occasional IMT may follow an aggressive clinical course. Rarely non-head and neck visceral IMTs have metastasized [466]. It is difficult to predict on the basis of histology which IMTs will be more aggressive [1156, 1363, 2842].
Lipoma

ICD-O code 8850/0

Lipomas of the larynx and hypopharynx comprise less than 0.5% of benign neoplasms at these sites, occur in all ages and affect both genders equally. The symptoms are non-specific but often include airway obstruction. In the larynx, supraglottic lesions predominate. Computed tomography and magnetic resonance document the lipomatous (low attenuation values and negative densitometry) nature and the extent of the mass. A lipoma is usually solitary, soft and sessile to polypoid. Tumours are composed of mature adipose cells, occasionally with foci of myxoid stroma. Distinction from well-differentiated liposarcoma is important. Association with systemic lipomatosis has been reported [1770]. Simple but complete excision is curative [329,409,1528,2756,2765].

Rhabdomyoma

Definition
A benign mesenchymal tumour with skeletal muscle differentiation and a propensity for occurrence in the head and neck.

ICD-O code 8900/0

Epidemiology
Based on histology rather than age, rhabdomyomas are divided into three types: fetal, juvenile (intermediate), and adult. Fetal rhabdomyomas (FRM) are 2-3 times more common in males and have been described in patients from birth to 65 years of age; about half of all patients are 15 years old or older at the time of diagnosis [597,1271]. Juvenile rhabdomyomas (JRM) are 2 times more common in males and have been observed in patients from 5 months to 58 years of age (average 18 years) [508,1271]. Adult rhabdomyomas (ARM) are 3-5 times more common in males and occur in an older population. About 80-90% of patients are over the age of 40 years (median 55-60 years, range 15 months to 82 years) [597,1272].

Etiology
Whether RMs are hamartomas or true neoplasms is controversial. Cytogenetics examination of an ARM has demonstrated clonal chromosomal abnormalities which supports a neoplastic origin [886]. Extracardiac RMs should be distinguished from those in the heart. Cardiac RMs are hamartomas and often associated with tuberous sclerosis. Extracardiac RMs, with the rare exception of a FRM

Leiomyoma

ICD-O code 8890/0

Leiomyomas (angioleiomyoma) of the larynx comprise less than 0.2% of all laryngeal neoplasms. The tumour has been reported in all age groups, but primarily in adults, and is somewhat more common in males. The ventricle and false vocal cord are sites of predilection.

Fig. 3.55 Rhabdomyoma of the hypopharynx in an adult. Note the characteristic tan colour and multinodularity.

Fig. 3.56 Rhabdomyoma. A Adult rhabdomyoma composed of large epithelioid cells with pink cytoplasm and eccentric nuclei with prominent nucleoli. The cytoplasmic vacuoles represent glycogen. B Diffuse desmin staining of cells and cross-strations in the cell in the upper middle portion of the illustration. C Juvenile (intermediate) rhabdomyoma showing spindle-shaped rhabdomyoblasts. D Fetal myxoid rhabdomyoma. Note the sharp border.
occurring in a few patients with the nevoid basal cell carcinoma syndrome, are virtually never associated with a phakomatosis [929].

**Localization**
RM of the larynx and hypopharynx are uncommon [1225]. In the larynx, they tend to centre around the true and false vocal cords and ventricles.

**Clinical features**
The tumours generally present as hoarseness, airway obstruction, dysphagia, or sensation of a foreign body in the throat.

**Macroscopy**
FRMs are usually 1-5 cm, circumscribed and grey-white to tan-pink with a mucoid cut surface.
ARMs are circumscribed, tan to red-brown and multinodular. Most are less than 5 cm, but may be larger.

**Histopathology**
FRMs vary from sparse to moderately cellular and are composed of immature cells with little cytoplasm and small, round to oval nuclei, sometimes with prominent nucleoli. The cytoplasm contains abundant glycogen, often appearing as vacuoles and producing a characteristic “spider cell”. Rod-shaped cytoplasmic crystals (“jackstraws”) may also be apparent. Cross striations are infrequent and, mitoses and necrosis are absent.
JRMs contain a large number of strap-shaped muscle cells with abundant eosinophilic cytoplasm with centrally located nuclei. Cytoplasmic vacuoles are common. The tumour often co-exists with typical areas of FRM.

**Immunoprofile**
RM are positive for muscle specific actin, smooth muscle actin, desmin, and myoglobin. ARMs are also variably weakly positive for vimentin (35% of cases) and even S-100 protein (67% of cases), but negative for glial fibrillary acidic protein (GFAP), cytokeratin, and epithelial membrane antigen. FRMs are also variably positive for vimentin (75% of cases), S-100 protein (50% of cases), and GFAP (50% of cases).

**Differential diagnosis**
The FRM must be distinguished from embryonal rhabdomyosarcoma. The lack of significant infiltration of adjacent tissues; absence of cellular pleomorphism, mitoses, and necrosis; and the presence of muscle maturation at the periphery of the lesion are features indicative of a rhabdomyoma.

ARMs are usually easily recognized. Infrequently, it may be confused with a pleomorphic rhabdomyosarcoma, granular cell tumour, hibernoma, oncocytoma, alveolar soft parts sarcoma, or crystal-storing histiocytosis [1268].

**Prognosis and predictive factors**
In contrast to FRMs, which are generally solitary lesions, 3-10% of ARMs may be multifocal, either synchronous or asynchronous. Conservative but complete excision is the treatment of choice. Ten to 40% of ARMs may recur, either within a few months or 10-15 years later. FRMs, in contrast, demonstrate less tendency for recurrence. Some of the “recurrent” ARMs may represent additional primary tumours in a patient with multifocal disease. RM have no malignant potential.

**Haemangioma and Lymphangioma**

**ICD-O-codes**
- Haemangioma: 9120/0
- Lymphangioma: 9170/0

Haemangiomas of the larynx are divided into juvenile (congenital) and adult types based on age of presentation, histologic appearance, and possibly patient outcome. Pediatric patients present at or within several months of birth with subglottic lesions that may result in potentially life-threatening airway obstruction and haemorrhage. In addition, about half of all pediatric patients with subglottic haemangiomas may have haemangiomas in other locations, most of which are cutaneous, rarely visceral. Adult haemangiomas are more often found in the supraglottic larynx. Grossly, haemangiomas are soft and compressible and range from red to blue, depending on the degree of vascularity. They may be either flat and diffuse or bulging and polypoid. The term ‘haemangiomatosis’ is sometimes used when the lesion is widespread and involves contiguous or non-contiguous sites. Microscopically, haemangiomas are categorized into capillary and cavernous types, and often demonstrate a lobular pattern of growth. Juvenile haemangiomas are usually cellular and of the capillary type while in adults, they are more often cavernous. Haemangiomas should be distinguished...
from telangiectasia, vascular stage of vocal cord polyps and granulation tissue. The distinction between haemangioma and telangiectasia may be difficult. But in the correct clinical setting of a positive family history of hereditary haemorrhagic telangiectasia (Rendu-Osler-Weber syndrome), typical lesions (in any location) and episodic bleeding can help to define the syndrome. Vascular vocal cord polyps occur exclusively on the true vocal cord and are separated from haemangiomas by a large amount of extravascular fibrin. Fibrin, if seen in haemangiomas, is always intravascular. The lobular growth of haemangioma distinguishes it from granulation tissue. If the lesion is biopsied rather than excised, unusually excessive bleeding may give a clue as to the type of lesion encountered. Although the preferred treatment is laser excision, therapy has included expectant management, systemic steroids, intralesional sclerosing agents and surgical excision [277,439,1235,1487].

**Granular cell tumour**

**Definition**
A neural tumour composed of round and/or spindle cells with pink, granular cytoplasm due to abundant intracytoplasmic lysosomes.

**ICD-O code** 9580/0

**Synonyms**
Granular cell myoblastoma, granular cell schwannoma, Abrikossoff tumour.

**Epidemiology**
Granular cell tumours (GCT) of the larynx affect both sexes equally and occur over a broad age range (4-70 years), with a mean of 34 years [2247]. They are uncommon in children. Only 20 cases were identified in 1998 in patients less than 17 years of age [1114].

GCTs of the trachea are even more unusual. Of 30 cases, 84% occurred in women and the peak incidence was the fourth decade (range 6-56 years) [315].

**Localization**
GCTs may occur anywhere in the larynx. In adults, the most common site is the posterior half of the true vocal cord, while in children the subglottis is the site of predilection. Tracheal tumours, in turn, arise most often in the cervical trachea.

**Clinical features**
GCTs are especially common in the Black population. Most patients have only a single tumour, but in 2-10% of individuals, multiple tumours may be found [924]. Hoarseness is the usual presenting symptom of laryngeal tumours while stridor or airway obstruction (often mistaken for asthma) is characteristic of those in the trachea.

**Macroscopy**
The majority of tumours are firm, polypoid or sessile, and less than two centimetres. They are covered by an intact (rarely ulcerated) mucosa, and on cross section are grey-white or yellow.
Histopathology
GCTs are poorly circumscribed and composed of round or spindle cells, often in a syncytial pattern. The nuclei are small, hyperchromatic, and centrally located. The cytoplasm is eosinophilic and contains numerous periodic acid-Schiff-positive, diastase-resistant granules. The granules are also S100 protein and CD68 positive, and ultrastructurally represent lysosomes.

Malignant granular cell tumour
Approximately 1-2% of all GCTs are malignant and exhibit either aggressive local behaviour or distant metastasis (lung, bone) [253,713]. Criteria for malignancy include: 1) necrosis, 2) spindling of cells, 3) vesicular nuclei with large nucleoli, 4) greater than 2 mitoses per 10 high power fields at 200X magnification, 5) high nuclear to cytoplasmic ratio, and 6) pleomorphism. Neoplasms that meet 3 or more of these criteria are classified as malignant. Those that meet only one or two criteria are regarded as atypical while those that show only focal pleomorphism but none of the other features are classified as benign.

Differential diagnosis
Small biopsies with sparse granular cells associated with pseudoepitheliomatous hyperplasia can easily be mistaken for squamous cell carcinoma. In the larynx, GCTs are typically non-ulcerated and occur on the posterior half of the true vocal cord in patients less than 50 years of age. In contrast, squamous cell carcinomas of the larynx are often ulcerated and arise on the anterior half of the true vocal cord in patients over the age of 50 years.

Prognosis and predictive factors
GCTs are radioresistant. Most can be removed endoscopically. Larger lesions may require an open excision. Although initial therapy is usually curative, 2-8% of patients may develop local recurrences. Some “recurrences”, however, may represent new primary lesions in a patient with multifocal disease.

Haematolymphoid tumours

Non-Hodgkin lymphoma
Primary non-Hodgkin lymphomas (NHL) of the hypopharynx, larynx or trachea are very rare. They account for 1% of all primary extranodal NHL [809]. By contrast, secondary laryngeal lymphomas are more common and represent spread from cervical and mediastinal lymph nodes, and thyroid gland. Patients present with hoarseness, foreign body sensation, or mild airway obstruction. Supraglottic tumours are more frequent, but all regions of the larynx can be involved.

Most primary laryngeal NHL are B-cell lymphomas, especially diffuse large B-cell lymphoma (DLBCL) and extranodal marginal zone B-cell lymphoma of MALT type [63,601,1285,1771]. Rare cases of extranodal NK/T cell lymphoma of nasal type [371,1761] and peripheral T-cell lymphoma [1285,1632,1761] have also been reported. Most patients (>90%) present with low clinical stage (Stage IE/IIE) [63,1285,1771], but occasional patients can succumb to acute laryngeal obstruction [1771]. NK/T cell lymphomas and peripheral T-cell lymphomas have a poorer outcome as compared to B-cell lymphomas [63,1285,1761,1771].

Patients with primary tracheal NHL may present with airway obstruction, dyspnoea, wheezing or cough. Most reported cases are extranodal marginal zone B-cell lymphoma of MALT type [762,1275]. Primary hypopharyngeal NHL is extremely rare [809]. Both extranodal NK/T cell lymphoma of nasal type [2455,2605] and extranodal marginal zone B-cell lymphoma of MALT type have been reported [2773].

Plasmacytoma
Definition
Plasmacytoma is a monoclonal plasma- cytotic proliferation. A soft tissue plasmacytoma without bone marrow involvement is referred to as extramedullary plasmacytoma (EMP).

ICD-O code 9734/3

Localization
The larynx and pharynx are the most common head and neck sites for extramedullary plasmacytoma. For details see Chapter 1 on sinonasal tumours (pp. 61-63).

A.C.L. Chan
J.K.C. Chan
S.B. Kapadia
Chondrosarcoma

Definition
Chondrosarcoma is a malignant tumour of the laryngeal framework characterized by the formation of neoplastic hyaline cartilage.

ICD-O code 9220/3

Epidemiology
Laryngeal chondrosarcoma (LCS) is the most common non-epithelial malignancy of the larynx, and comprises 75% of laryngeal sarcomas (1489). Cartilaginous neoplasms are estimated to represent 0.07-0.2% of all laryngeal tumours (255). Approximately 300 cases of cartilaginous laryngeal tumours have been reported; the majority are represented by chondrosarcomas. They affect adults in the 6-9th decades, with a mean age at diagnosis of 60-65 years. The male to female ratio is approximately 3:1.

Localization
LCS arise predominantly in the ossified hyaline cartilages. The cricoid ring is the most frequently involved, especially its posterior or posterolateral aspect, followed by the thyroid cartilage. Bulky tumours may encompass both structures, obscuring the exact site of origin. Chondrosarcoma of the epiglottis has only been rarely reported.

Clinical features
Patients with LCS typically present with hoarseness, and/or airway obstruction and dyspnoea. An external neck mass may be noted when it arises in the thyroid lamina. These tumours grow slowly and can be asymptomatic until they reach considerable size. On examination, the usual appearance is that of a subglottic swelling with intact mucosa. Vocal cord paralysis is a common finding at presentation. LCS have characteristic features on CT examination, with expansion of the affected cartilage by a relatively circumscribed, hypodense mass containing stippled to coarse calcifications. MR imaging may show better definition of the tumour boundaries but is less likely to detect internal calcifications (255,1888, 2598). LCS are notoriously difficult to biopsy. Their characteristic imaging appearance precludes the necessity for preoperative biopsy.

Macroscopy
LCS are bulky, lobulated neoplasms that expand and distort the involved site. The cut surface is firm to hard, translucent, pale grey-blue, with gritty calcifications. Myxoid change, if present, is characterized by soft, cystic, or gelatinous areas. Dedifferentiated chondrosarcoma reveals fleshy areas resembling high-grade sarcoma.

Histopathology
The diagnosis and grading of LCS is based on general criteria for chondrosarcoma (1509). The low power architecture of LCS is that of a lobulated neoplasm with pushing borders. The tumour periphery shows invasive growth of neoplastic lobules into adjacent soft tissue or the marrow spaces of ossified cartilage. The overwhelming majority of LCS are low or intermediate grade, with variability from area to area (28,255,1152,1888).
Tumours of bone and cartilage

High-grade LCS are generally considered rare; although in the largest reported series they comprised 5% [2598]. Metaplastic bone formation and calcification are also common to LCS. Myxoid change is infrequent. Dedifferentiated and clear cell variants of LCS have been reported, albeit rarely [255, 1888, 2598]. Dedifferentiated chondrosarcoma is characterized by the presence of two distinct components: well-differentiated chondrosarcoma and a high-grade, non-cartilaginous sarcoma. Clear cell chondrosarcoma is composed of chondrocytes with abundant clear cytoplasm and prominent cell membranes. LCS, similar to chondrosarcoma found elsewhere, expresses S-100 protein (strongly) and vimentin (focally) in immunohistochemical studies [255, 1888].

Differential diagnosis

Low-grade LCS may be difficult to distinguish from chondroma. Grade 1 LCS show subtle increases in cellularity, with nuclear hyperchromasia and occasional binucleate forms. Irregular clustering of cell groups, or “cluster disarray” is found in all grades of LCS [1152] and may be a useful feature in the distinction of low-grade chondrosarcoma from chondroma. The diagnosis of laryngeal chondroma should be reserved for small (less than 1-2 cm), clinically insignificant lesions, without discernible atypia. Any recurrent cartilaginous tumour of the larynx should be considered a chondrosarcoma [255]. Chondrometaplasia is characterized by small (less than 1 cm) nodules of bland, fibroelastic cartilage which are found in the submucosal soft tissue of the glottic region [747,1161].

Precursor lesions

Understanding of the biologic potential of laryngeal cartilaginous neoplasms has evolved in recent years such that many reported laryngeal chondromas would now be interpreted as low-grade LCS [1888,2178,2598]. (See section on Chondromas).

Histogenesis

LCS usually develop in ossified cartilage. A pluripotential mesenchymal stem cell, which may be recruited in the process of ossification, is postulated to be the cell of origin [255]. Several authors have proposed that LCS may develop in a benign chondroma [2598]; this hypothesis remains highly controversial.

Prognosis and predictive factors

LCS are more indolent than chondrosarcomas arising elsewhere [2598]. This may be due to the fact that LCS are symptomatic at a smaller size when compared to their skeletal counterparts. LCS are managed with conservative surgery [255,1489,2598]. Incomplete resection (shelling out) is associated with local recurrence [2178]. Metastases from LCS, usually pulmonary, are distinctly unusual (<10%), and related to higher grade or dedifferentiation [2598]. LCS related mortality is very low [1489]. Myxoid change involving greater than 10% of the neoplasm correlates adversely with outcome [2598].

Osteosarcoma

Definition

Osteosarcoma is a malignant tumour characterized by the direct formation of osteoid by neoplastic cells.
Tumours of the hypopharynx, larynx and trachea

ICD-O code 9180/3

Epidemiology
Laryngeal osteosarcoma (LOS) is extremely rare, with fewer than 20 documented cases [198, 526, 926, 1604, 1807, 2054, 2154, 2203, 2327, 2675]. LOS affects an older age group than osteosarcoma arising in long bone [2054]. They manifest in patients from about 50-80 years of age, nearly exclusively in males.

Localization
These sarcomas usually arise from the endolaryngeal soft tissue, vocal cords and/or anterior commissure, rather than the laryngeal framework.

Clinical features
LOS is typically a polypoid mass that impinges on the airway. Symptoms depend on the tumour site; hoarseness, dyspnoea, and airway obstruction are the most frequent complaints. Imaging studies reveal an invasive, mineralized mass, either situated primarily in the soft tissue of the glottis or expanding one of the laryngeal cartilages [1807, 2203]. Biopsy may be difficult when the tumour is heavily mineralized.

Macroscopy
These sarcomas usually arise from the endolaryngeal soft tissue, vocal cords and/or anterior commissure, rather than the laryngeal framework.

Clinical features
LOS is typically a polypoid mass that impinges on the airway. Symptoms depend on the tumour site; hoarseness, dyspnoea, and airway obstruction are the most frequent complaints. Imaging studies reveal an invasive, mineralized mass, either situated primarily in the soft tissue of the glottis or expanding one of the laryngeal cartilages [1807, 2203]. Biopsy may be difficult when the tumour is heavily mineralized.

Histopathology
Similar to other primary mesenchymal neoplasms, LOS typically retains a “Grenz zone” of tumour-free, superficial submucosa just beneath the non-dysplastic epithelium. LOS is uniformly high-grade [1604, 2054], composed of pleomorphic spindle cells. The amount of osteoid produced by the malignant cells can vary, but characteristic lace-like osteoid is present at least focally [926, 1604]. Tumour giant cells may also be identified [198, 526, 1807]. Laryngeal osteosarcoma may also represent the high-grade sarcomatous component of dedifferentiated chondrosarcoma of the larynx [2675].

Differential diagnosis
LOS must be distinguished from spindle cell carcinoma (SPCC) with heterologous osteoid production, since the spindle cell component of the latter tumour may be highly pleomorphic and focally produce osteoid [1604]. In contrast to LOS, SPCC is characterized by either 1) a concomitant squamous abnormality (dysplasia, carcinoma in-situ, or invasive SCC) or 2) evidence of epithelial differentiation in spindle cells. In addition, the malignant spindle cells of SPCC usually abut the overlying mucosa or merge with its basal layer in a feathering pattern, without the presence of a Grenz zone.

Prognosis and predictive factors
Reported cases of LOS have demonstrated aggressive clinical behaviour. Local recurrence is frequent, as are distant metastases, typically involving the lung. At least half of the reported patients have died of disease, most within a year of diagnosis [198, 1604].

Chondroma

Definition
A benign tumour composed of mature hyaline cartilage

ICD-O code 9220/0

Epidemiology
True chondromas of the larynx are extremely unusual and are greatly outnumbered by laryngeal chondrosarcoma (LCS) [2593]. The age incidence is difficult to estimate as many previous reports of chondroma probably represent LCS [465, 591].

Clinical features
Chondromas of the larynx primarily affect the cricoid and thyroid cartilages. Chondroma may be an incidental finding, or cause minor symptoms such as hoarseness. A clinically significant cartilaginous tumour is more likely to be LCS than a chondroma.

Macroscopy
Chondromas are well-circumscribed tumours. The cut surface is uniform, with a translucent, pale grey-blue appearance. A cartilaginous neoplasm greater than 2 cm in dimension more likely represents LCS.

Histopathology
Chondromas are composed of benign chondrocytes producing hyaline cartilage. They may show a lobular growth pattern. The appearance is uniform and monotonous with overall low cellularity. The chondrocytes are relatively evenly distributed, lack nuclear pleomorphism and mitotic activity and contain a single nucleus per lacuna [591, 1489]. In LCS, the neoplastic chondrocytes are distributed in cell groups of varying size and cellularity (cluster disarray). However, because the histopathology of LCS is variable within a tumour, thorough sampling of any cartilaginous tumour is recommended. The diagnosis of chondroma should be reserved for small lesions that have been completely excised and entirely examined [591].

Prognosis and predictive factors
Laryngeal chondroma does not recur after conservative excision. Any recurrent cartilaginous neoplasm of the larynx should be interpreted as LCS [591, 2593].
**Giant cell tumour**

**Definition**
A benign but locally destructive neoplasm composed of sheets of ovoid to spindle-shaped mononuclear cells with uniformly dispersed osteoclast-like giant cells.

**ICD-O code**
9250/1

**Synonym**
Osteoclastoma

**Epidemiology**
Giant cell tumours of the larynx are very rare (590,982,1092,1642,2163). They represented only 0.09% of almost 9000 benign and malignant laryngeal tumours (2785). In total 28 cases have been reported comprised of 25 men and 3 women, aged 23-62 years (mean about 40-45 years).

**Localization**
The thyroid cartilage is most commonly involved, followed by cricoid cartilage and epiglottis (2785).

**Clinical features**
The tumours enlarge slowly and manifest as palpable neck masses, hoarseness, airway obstruction, dysphagia or sore throat. On imaging, it often appears as a tumour exploding from within the cartilage, destroying it and extending into soft tissue of the neck or endolarynx.

**Macroscopy**
Most tumours have ranged from 2.4-7cm (mean about 4.4-5cm) and have been centred in the thyroid or cricoid cartilages, especially in the normally ossified portions of these cartilages (2785). On cut section, they are soft, red to grey-pink and frequently extend beyond the cartilage into the adjacent soft tissue. Haemorrhage and cystic degeneration are common.

**Histopathology**
The tumours are similar histologically to the giant cell tumour of bone and, as such, consist of a dual population of cells: mononuclear cells and osteoclast-like giant cells. The mononuclear cells appear as broad sheets of cells reminiscent of histiocytes. They are round, ovoid or spindled and have pink to amphiphilic cytoplasm and round, vesicular nuclei with occasional prominent nucleoli. The giant cells are evenly distributed throughout the tumour and contain up to 20 or more nuclei per cell. The nuclei of the giant cells are identical to those of the mononuclear cells.

The stroma is vascular and contains many thin-walled vessels with small areas of haemorrhage and haemosiderin-laden macrophages. Mitoses are usually seen, averaging 4 per 10 high power fields (range 1-12 per 10 high power fields) (2785). Atypical mitoses are not seen.

**Differential diagnosis**
This includes giant cell (reparative) granuloma, brown tumour of hyperparathyroidism, fibrous histiocytoma and a pleomorphic carcinoma. Giant cell granuloma of the cricoid cartilage is exceptionally rare (2587). The giant cells in this tumour are not evenly distributed, but rather concentrate around areas of recent and/or old haemorrhage. A giant cell granuloma also exhibits more stromal fibrosis. A brown tumour is identical histologically to the giant cell granuloma, but is associated with elevated serum calcium. A benign fibrous histiocytoma contains a more uniform storiform arrangement of fibroblasts and does not show a symmetrical distribution of giant cells. A malignant fibrous histiocytoma (giant cell type) exhibits significant nuclear pleomorphism and abnormal mitoses, none of which are seen in a giant cell tumour. A pleomorphic carcinoma will not only show abnormal mitoses and pleomorphism, but will also be positive for cytokeratin.

**Prognosis and predictive factors**
Complete, but conservative surgical excision is the treatment of choice. Large tumours may require a partial or total laryngectomy. Adjuvant therapy is unnecessary. There have been no convincing records of local recurrence or malignant behaviour secondary to giant cell tumour of the laryngeal framework (1138,1181).
Definition
Primary laryngeal mucosal malignant melanomas (PLMMM) are neural crest-derived neoplasms originating from melanocytes and demonstrating melanocytic differentiation.

ICD-O code 8720/3

Epidemiology
Approximately 15-20% of all malignant melanomas arise in head and neck sites, and of these over 80% are of cutaneous origin. Of the approximate remaining 20%, the majority are of ocular origin; mucosal malignant melanomas (MMM) of the upper aerodigestive tract represent from 0.5-3% of melanomas of all sites [735]. In the upper aerodigestive tract, the most common site of occurrence is the sinonasal tract. PLMMM are extremely rare, with less than 60 cases reported in the literature [51,77,831,1277,1516,1668,1843,2754,2755]. PLMMM are much more common in men than in women with over 80% of cases occurring in men [2754]. PLMMM occur over a wide age range from 35-86 years with an average age of 58 years, and are most frequent in the 6th and 7th decades of life. Most cases of PLMMM occur in Caucasians but Blacks are also affected.

Etiology
There are no known etiologic factors for PLMMM. Melanosis [912,2022], intralaryngeal naevi [2263,2287] and lentigo [2630] of the larynx have been reported. It has been suggested that PLMMM may arise from malignant degeneration of intralaryngeal melanocytes or melanocytic lesions [2754].

Localization
The majority (more than 60%) of PLMMM occur in the supraglottic larynx [2754,2755], including the epiglottis, arytenoids, aryepiglottic folds, ventricle, false vocal cord, and pyriform sinus [2755]. Other less common sites of occurrence include the glottic region along the true vocal cord and the posterior commissure. To date, there are no documented reports of PLMMM involving the subglottic region.

Clinical features
The clinical presentation of PLMMM includes hoarseness, dysphagia, sore throat, intermittent haemoptysis, neck or jaw pain and a cervical neck mass. Symptoms generally occur over short periods of time, ranging from 3-6 months [2755]. Multicentric (synchronous, metachronous) MMM of other upper aerodigestive tract sites are not typically present.

Macroscopy
The macroscopic appearance of PLMMM vary and include nodular, mul-
berry-like, sessile, polypoid, exophytic or pedunculated lesions with equally variable colour, including black, brown, red-pink, tan-grey and white (2754). The size of the tumours range from 3-4 mm up to 8.0 cm in greatest dimension (2754).

**Histopathology**

PLMMM are identical to melanomas at other sites. In the presence of an intact laryngeal mucosa, continuity of the tumour with the surface epithelium (i.e., junctional or pagetoid changes) can be identified; however, even in the presence of intact surface epithelium, junctional changes may not be seen. Given the fact that normal melanocytes may localize to the submucosal compartment within minor mucoserous glands or within the stroma (2754, 2755), junctional change is not required to render a diagnosis of PLMMM.

**Prognosis and predictive factors**

PLMMM has a poor prognosis. The average survival rate is usually less than 3.5 years (2159, 2754) with a 5-year survival rate of less than 20% (2754). Radical surgical excision is the treatment of choice. Adjuvant radiotherapy and chemotherapy are of questionable value in the management of PLMMM. Approximately 80% of patients with PLMMM have metastatic disease to the regional lymph nodes as well as to distant viscera (e.g., brain, lungs, bone). Pathologic criteria that are used to predict the biologic behaviour in association with cutaneous melanomas, including the depth of invasion, age and gender of the patient, and cytomorphology generally do not apply for PLMMM (2754, 2755). Further, prognostic significance has not been found for tumour thickness, level of invasion, ulceration, mitotic index or nerve/nerve sheath involvement for PLMMM (2080).

**Fig. 3.70** Primary laryngeal mucosal malignant melanoma. Prominent and obvious intracytoplasmic melanin deposition is seen in this PLMMM; this extent of melanin deposition is unusual for MMM.

**Fig. 3.71** Primary laryngeal mucosal malignant melanoma. Immunohistochemical reactivity in this epithelioid melanoma includes: A HMB-4 B S100 protein. C melan A and D vimentin.
Tumours of the hypopharynx, larynx and trachea

Definition
Tumours involving the hypopharynx, larynx and/or trachea that originate from, but are not in continuity with, other primary malignant neoplasms. Leukemias and lymphomas are excluded.

Epidemiology
Metastases to the larynx are uncommon. Only 11 cases over 20 years were identified in one series [160]. Eight additional cases were found in a review of more than 4000 laryngeal malignancies [7353]. In 1993, 134 cases were recorded in the world literature [735]. Metastases to the hypopharynx and trachea are even more unusual.

Age and sex distribution
Laryngeal metastases increase with age (median 58 years, range 24-83 years) and are more common in males by a ratio of 2:1 [160,741].

Etiology
The overwhelming majority of tumours that metastasize to the larynx are either malignant melanomas or carcinomas. Only 5% or fewer are from mesenchymal tumours (bone and soft tissue sarcomas).

Table 3.7 Site of origin of 120 tumours metastatic to larynx*

<table>
<thead>
<tr>
<th>Site</th>
<th>Frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin (melanoma)</td>
<td>39.1%</td>
</tr>
<tr>
<td>Kidney</td>
<td>13.3%</td>
</tr>
<tr>
<td>Breast</td>
<td>9.2%</td>
</tr>
<tr>
<td>Lung</td>
<td>7.5%</td>
</tr>
<tr>
<td>Prostate</td>
<td>6.7%</td>
</tr>
<tr>
<td>Colon</td>
<td>3.3%</td>
</tr>
<tr>
<td>Stomach</td>
<td>2.5%</td>
</tr>
<tr>
<td>Miscellaneous</td>
<td>18.4%</td>
</tr>
</tbody>
</table>

*Data based on reference [741]

Table 3.8 Site of origin of 12 tumours metastatic to trachea*

<table>
<thead>
<tr>
<th>Site</th>
<th>Frequency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breast</td>
<td>33%</td>
</tr>
<tr>
<td>Colon</td>
<td>25%</td>
</tr>
<tr>
<td>Skin (melanoma)</td>
<td>17%</td>
</tr>
<tr>
<td>Uterine cervix</td>
<td>8%</td>
</tr>
<tr>
<td>Endometrium</td>
<td>8%</td>
</tr>
<tr>
<td>Ovary</td>
<td>8%</td>
</tr>
</tbody>
</table>

*Data based on references [179,485,1365,1599,1889,2775]

Fig. 3.72. Adenocarcinoma of the colon metastatic to the larynx. Note the involvement of both soft tissue and the inferior border of the thyroid cartilage.

Localization
Metastases to the larynx may be submucosal, cartilaginous or both. If cartilage is involved, it is usually only in the portion which has undergone ossification.

Clinical features
Generally, metastatic tumours to the larynx present with the usual supraglottic or glottic symptomatology. Richly vascular tumours, such as renal cell carcinomas and thyroid carcinomas, often result in haemoptysis. On rare occasions, the metastasis is the only evidence of an otherwise occult primary tumour.

Tumour spread and staging
The majority of metastases to the larynx are haematogenous through the systemic circulation or the paravertebral venous plexus.

Prognosis and predictive factors
Metastases to the larynx, trachea or hypopharynx are usually associated with terminal, widespread disseminated disease. In some instances, the metastasis may be isolated or localized and, with appropriate therapy, a prolonged survival can be achieved.
CHAPTER 4

Tumours of the Oral Cavity and Oropharynx

Squamous cell carcinomas amount to more than 90% of malignant tumours of the oral cavity and oropharynx. As in other parts of the upper aerodigestive tract, there is a strong and synergistic association with tobacco smoking and alcohol abuse. In some regions, particularly the Indian subcontinent, oral cancer is among the most frequent malignancies, largely due to tobacco chewing.

The WHO Working Group has made an attempt to unify the terminology used to define the histological features of precursor lesions throughout the head and neck region. Although there has been considerable progress in the understanding of the genetic and molecular events underlying the progression of precancerous lesions to invasive carcinomas, this has yet to be translated into novel therapeutic strategies.
**WHO classification of tumours of the oral cavity and oropharynx**

<table>
<thead>
<tr>
<th>Malignant epithelial tumours</th>
<th>WHO Classification</th>
<th>Morphology Code</th>
</tr>
</thead>
<tbody>
<tr>
<td>Squamous cell carcinoma</td>
<td>Myoepithelial carcinoma</td>
<td>8982/3</td>
</tr>
<tr>
<td>Verrucous carcinoma</td>
<td>Carcinoma ex pleomorphic adenoma</td>
<td>8941/3</td>
</tr>
<tr>
<td>Basaloid squamous cell carcinoma</td>
<td>Salivary gland adenomas</td>
<td>8940/0</td>
</tr>
<tr>
<td>Papillary squamous cell carcinoma</td>
<td>Pleomorphic adenoma</td>
<td>8940/0</td>
</tr>
<tr>
<td>Spindle cell carcinoma</td>
<td>Myoepithelioma</td>
<td>8982/0</td>
</tr>
<tr>
<td>Acantholytic squamous cell carcinoma</td>
<td>Basal cell adenoma</td>
<td>8147/0</td>
</tr>
<tr>
<td>Adenosquamous carcinoma</td>
<td>Canalicular adenoma</td>
<td>8149/0</td>
</tr>
<tr>
<td>Carcinoma cuniculatum</td>
<td>Duct papilloma</td>
<td>8503/0</td>
</tr>
<tr>
<td>Lymphoepithelial carcinoma</td>
<td>Cystadenoma</td>
<td>8440/0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Epithelial precursor lesions</th>
<th></th>
<th></th>
</tr>
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<tbody>
<tr>
<td>Benign epithelial tumours</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Papillomas</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Squamous cell papilloma and verruca vulgaris</td>
<td>8050/0</td>
<td></td>
</tr>
<tr>
<td>Condyloma acuminatum</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Focal epithelial hyperplasia</td>
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</tr>
<tr>
<td>Granular cell tumour</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Keratoacanthoma</td>
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<td></td>
</tr>
<tr>
<td>Salivary gland tumours</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Salivary gland carcinomas</td>
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<td></td>
</tr>
<tr>
<td>Acinic cell carcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adenoid cystic carcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polymorphous low-grade adenocarcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basal cell adenocarcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Epithelial-myoeoepithelial carcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Clear cell carcinoma, not otherwise specified</td>
<td>8500/3</td>
<td></td>
</tr>
<tr>
<td>Mucinous adenocarcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oncocytic carcinoma</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Salivary duct carcinoma</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

1. Morphology code of the International Classification of Diseases for Oncology (ICD-O) (821) and the Systematized Nomenclature of Medicine (http://snomed.org). Behaviour is coded /0 for benign tumours, /3 for malignant tumours, and /1 for borderline or uncertain behaviour.
## TNM classification of carcinomas of the lip and oral cavity

<table>
<thead>
<tr>
<th>T</th>
<th>Primary tumour</th>
</tr>
</thead>
<tbody>
<tr>
<td>TX</td>
<td>Primary tumour cannot be assessed</td>
</tr>
<tr>
<td>T0</td>
<td>No evidence of primary tumour</td>
</tr>
<tr>
<td>Tis</td>
<td>Carcinoma in situ</td>
</tr>
<tr>
<td>T1</td>
<td>Tumour 2 cm or less in greatest dimension</td>
</tr>
<tr>
<td>T2</td>
<td>Tumour more than 2 cm but not more than 4 cm in greatest dimension</td>
</tr>
<tr>
<td>T3</td>
<td>Tumour more than 4 cm in greatest dimension</td>
</tr>
<tr>
<td>T4a</td>
<td>Tumour invades through cortical bone, inferior alveolar nerve, floor of mouth, or skin (chin or nose)</td>
</tr>
<tr>
<td>T4a (lip)</td>
<td>Tumour invades through cortical bone, into deep/extrinsic muscle of tongue (genioglossus, hyoglossus, palatoglossus, and styloglossus), maxillary sinus, or skin of face</td>
</tr>
<tr>
<td>T4b (lip and oral cavity)</td>
<td>Tumour invades masticator space, pterygoid plates, or skull base; or encases internal carotid artery</td>
</tr>
</tbody>
</table>

**Note:** Superficial erosion alone of bone/tooth socket by gingival primary is not sufficient to classify a tumour as T4.

<table>
<thead>
<tr>
<th>N</th>
<th>Regional lymph nodes</th>
</tr>
</thead>
<tbody>
<tr>
<td>NX</td>
<td>Regional lymph nodes cannot be assessed</td>
</tr>
<tr>
<td>N0</td>
<td>No regional lymph node metastasis</td>
</tr>
<tr>
<td>N1</td>
<td>Metastasis in a single ipsilateral lymph node, 3 cm or less in greatest dimension</td>
</tr>
<tr>
<td>N2</td>
<td>Metastasis as specified in N2a, 2b, 2c below</td>
</tr>
<tr>
<td>N2a</td>
<td>Metastasis in a single ipsilateral lymph node, more than 3 cm but not more than 6 cm in greatest dimension</td>
</tr>
<tr>
<td>N2b</td>
<td>Metastasis in multiple ipsilateral lymph nodes, none more than 6 cm in greatest dimension</td>
</tr>
<tr>
<td>N2c</td>
<td>Metastasis in bilateral or contralateral lymph nodes, none more than 6 cm in greatest dimension</td>
</tr>
<tr>
<td>N3</td>
<td>Metastasis in a lymph node more than 6 cm in greatest dimension</td>
</tr>
</tbody>
</table>

**Note:** Midline nodes are considered ipsilateral nodes.

<table>
<thead>
<tr>
<th>M</th>
<th>Distant metastasis</th>
</tr>
</thead>
<tbody>
<tr>
<td>MX</td>
<td>Distant metastasis cannot be assessed</td>
</tr>
<tr>
<td>M0</td>
<td>No distant metastasis</td>
</tr>
<tr>
<td>M1</td>
<td>Distant metastasis</td>
</tr>
</tbody>
</table>

### Stage grouping

<table>
<thead>
<tr>
<th>Stage</th>
<th>Tis</th>
<th>N0</th>
<th>M0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stage I</td>
<td>T1</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage II</td>
<td>T2</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage III</td>
<td>T1, T2</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T3</td>
<td>N0, N1</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IVA</td>
<td>T1, T2, T3</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T4a</td>
<td>N0, N1, N2</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IVB</td>
<td>T4b</td>
<td>Any N</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>Any T</td>
<td>N3</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IVC</td>
<td>Any T</td>
<td>Any N</td>
<td>M1</td>
</tr>
</tbody>
</table>


## TNM classification of carcinomas of the oropharynx

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</tr>
<tr>
<td>T4a</td>
<td>Tumour invades any of the following: larynx, deep/extrinsic muscle of tongue (genioglossus, hyoglossus, palatoglossus, and styloglossus), maxillary pterygoid, hard palate, and mandible</td>
</tr>
<tr>
<td>T4b</td>
<td>Tumour invades any of the following: lateral pterygoid muscle, pterygoid plates, lateral nasopharynx, skull base; or encases the carotid artery</td>
</tr>
</tbody>
</table>

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<tr>
<th>Stage</th>
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<th>N0</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Stage 0</td>
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<td>T1</td>
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<td>M0</td>
</tr>
<tr>
<td>Stage II</td>
<td>T2</td>
<td>N0</td>
<td>M0</td>
</tr>
<tr>
<td>Stage III</td>
<td>T1, T2</td>
<td>N1</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T3</td>
<td>N0, N1</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IVA</td>
<td>T1, T2, T3</td>
<td>N2</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>T4a</td>
<td>N0, N1, N2</td>
<td>M0</td>
</tr>
<tr>
<td>Stage IVB</td>
<td>T4b</td>
<td>Any N</td>
<td>M0</td>
</tr>
<tr>
<td></td>
<td>Any T</td>
<td>N3</td>
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</tr>
<tr>
<td>Stage IVC</td>
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<td>Any N</td>
<td>M1</td>
</tr>
</tbody>
</table>

**Note:** The regional lymph nodes are the cervical nodes.

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1 (847,2418).
2 A help desk for specific questions about the TNM classification is available at www.uicc.org/index.php?id=508 .
Tumours of the oral cavity and oropharynx: Introduction

Tumours of the oral cavity and oropharynx may be either epithelial, mesenchymal, or haematolymphoid. The epithelial tumours may be classified as those originating within the epithelium lining of the oral cavity and oropharynx and those derived from salivary gland tissue. Both will be included in this chapter, including precursor lesions where appropriate.

For the haematolymphoid diseases, the reader is referred to the WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues [1197], for mesenchymal ones to the WHO Classification of Tumours of Soft Tissue and Bone [775].

Oral Cavity

The oral cavity extends from the lips to the palatoglossal folds. The outer vestibule is enclosed by the cheeks and lips and forms a slit-like space separating it from the gingivae and teeth. It is limited above and below by mucosal reflections from the lips and cheeks.

The space bordered by the teeth and gingivae is the oral cavity proper. It is bounded inferiorly by the floor of the mouth and tongue and superiorly by the hard palate. The buccal mucosa extends from the commissure of the lips anteriorly to the palatoglossal fold posteriorly. It is lined by thick, non-keratinized stratified squamous epithelium and contains variable numbers of sebaceous glands (Fordyce spots or granules) and minor salivary glands. The duct of the parotid gland (Stensen's duct) opens on a papilla or fold opposite the upper second permanent molar tooth.

The mucous membrane related to the teeth is the gingiva. The gingival mucosa surrounds the necks of the teeth and the alveolar mucosa overlies the alveolar bone and extends to the vestibular reflections. The junction between these two parts is marked by a faint scalloped line called the mucogingival junction. The gingival mucosa is pink and firmly attached to the underlying bone and necks of the teeth (attached gingiva) except for a free marginal area. It is usually non-keratinized or parakeratinized. The alveolar mucosa is reddish and covered by thin, non-keratinized stratified squamous epithelium. Minor salivary glands may be seen in the alveolar mucosa and occasionally the attached gingiva.

The hard palate is continuous anteriorly with the maxillary alveolar arches and posteriorly with the soft palate. A median raphe extends anteriorly from this junction to the incisive fossa into which the nasopalatine foramen opens. Most of the palatal mucosa is firmly bound to the underlying bone forming a mucoperiosteum. It is covered by orthokeratinized stratified squamous epithelium and posteriorly contains many minor mucous salivary glands.

The oral part of the tongue (anterior two thirds) lies in front of the V-shaped sulcus terminalis. It is mobile and attached to the floor of the mouth anteriorly by a median lingual frenum. The dorsal part is covered by stratified squamous epithelium and contains several types of papillae. The most numerous are the hair-like filiform papillae which are heavily keratinized. There are less numerous and evenly scattered fungiform papillae which form pink nodules and contain taste buds. Taste buds here and in other oral sites are occasionally mistaken for junctional melanocytic proliferation or Pagetoid infiltration. In front of the sulcus terminalis there are 10-12 circumvallate papillae. These contain many taste buds on the surface and in a deep groove that surrounds each papilla. In addition, the ducts of minor serous salivary glands (von Ebner's glands) open into the base of the groove. At the postero-lateral aspect of the tongue where it meets the palatoglossal fold there are the leaf shaped foliate papillae. These also may contain taste buds on the surface and the core of the papillae often contains lymphoid aggregates similar to those in the rest of the Waldeyer ring. In addition, there are minor salivary glands in the underlying lingual musculature. The ventrum of the tongue is covered by thin, non-keratinized stratified squamous epithelium which is continuous with similar mucosa in the floor of the mouth. Minor salivary glands (glands of Blandin and Nuhn) are present, predominantly towards the midline and deep within the lingual musculature. They can extend to involve the tip of the tongue.

The floor of the mouth is a horseshoe-shaped area between the ventrum of the tongue medially and the gingivae of the lower teeth anteriorly and laterally. It extends to the palatoglossal folds distally and is in continuity with the retromolar pad behind the lower third molar tooth. The mucosa covers the major sublingual glands and the submandibular (Wharton's) ducts which open anteriorly onto the submandibular papillae on either side of the median sublingual frenum. It is important to note that 75% of oral squamous cell carcinomas have been reported to arise in an area that comprises the floor of the mouth and adjacent lingual mucosa, sublingual sulcus and retromolar region [1767]. This region forms only about 20% of the total mucosal area. The zone of increased susceptibility has been called the ‘drainage area’ as it is thought that any carcinogens present in the mouth pool there before being swallowed. It is obvious, therefore, that any precursor lesions in these areas should be regarded as highly suspicious.

Oropharynx

The oropharynx lies behind the oral cavity. It is bounded superiorly by the soft palate and inferiorly by a hypothetical...
horizontal line level with the tip of the epiglottis. Anteriorly are the isthmus of the fauces and the posterior third of the tongue, and the lateral wall is formed by the palatopharyngeal arches and the palatine tonsils. The posterior wall contains the pharyngeal tonsils. The palatine tonsils are two masses of lymphoid tissue situated in the triangular recess (tonsillar sulcus) between the anterior and posterior faucial pillars. They extend from the soft palate to the dorsum of the tongue. The surface is convoluted and deep clefts or crypts can penetrate almost its full thickness. The bulk of the tonsil consists of lymphoid tissue arranged in nodules or follicles. There are no afferent lymphatics and no subcapsular sinuses. Squamous cell carcinomas at this site can invade deeply into the underlying tissues, base of tongue and lateral pharyngeal wall. They also have a particular tendency to extend upwards into the nasopharynx. The soft palate is a mobile, muscular flap attached to the posterior edge of the hard palate and extending to a free margin posteriorly. The uvula forms a small, conical, midline process. The oral surface of the soft palate is covered by non-keratinized stratified squamous epithelium and contains many minor mucous glands. The uvula contains mainly fat and a few muscle fibres but minor salivary glands may also be seen and occasionally salivary gland tumours develop at this site. The pharyngeal part of the tongue is immobile and has a bossellated surface due to the presence of underlying lymphoid tissue forming the lingual tonsils. Minor salivary glands are also present.

**Lymphatic drainage of mouth and oropharynx**

The main sites of lymphatic drainage from the mouth and oropharynx are the jugulodigastric, submandibular and submental lymph nodes. Lymph vessels from the gingiva usually drain to the submandibular lymph nodes but those in the lower incisor region run to the submental nodes. Most of the vessels from the palate run to the jugulodigastric group but some involve the retropharyngeal nodes. There is a rich lymphatic plexus in the tongue and the main vessels can be subdivided into marginal and central. The marginal vessels drain the lateral third of the dorsum and contiguous lateral border and part of the ventrum of the tongue. They run to the ipsilateral submandibular nodes. Those towards the tip of the tongue drain to the submental nodes. Central lymph vessels drain to the submandibular nodes on both sides. Some marginal and central vessels run directly to the jugulodigastric group but some can pass direct to the jugulo-omohyoid nodes. Vessels from the area of the circumvallate papillae and posterior third of the tongue drain to the jugulodigastric, jugulo-omohyoid or intermediate nodes, either unilaterally or bilaterally. Most of the lymphatics of the palatine tonsils drain to the jugulodigastric nodes.
Squamous cell carcinoma

Definition
An invasive epithelial neoplasm with varying degrees of squamous differentiation and a propensity to early and extensive lymph node metastases, occurring predominantly in alcohol and tobacco-using adults in the 5th and 6th decades of life.

ICD-O code 8070/3

Epidemiology
More than 90% of malignant neoplasms of the oral cavity and oropharynx are squamous cell carcinomas of the lining mucosae with relatively rare neoplasms arising in minor salivary glands and soft tissues. It is important to specify which anatomical sites are included in epidemiological data. Separate assessment of incidence rates for the oral cavity and oropharynx is complicated by the difficulty of assigning a site of origin to tumours that are often advanced.

Males are affected more often than females because of heavier indulgence in both tobacco and alcohol habits in most countries: in India the highest rates of intraoral cancer may be found in women who chew tobacco heavily. The male to female ratio is, however, globally lower for cancer of the oral cavity than for cancer of the oropharynx, perhaps suggesting that higher exposure to tobacco smoking and alcohol drinking are required to induce oropharyngeal than oral cancer (796). Globally some 389,650 cases occurred in the year 2000; 266,672 for the oral cavity (ICD-9 140-5) and 122,978 for the oropharynx (ICD-9 146,8-9) (1981). This represents 5% of all cancers for men and 2% for women.

In males, the country with the highest rate in the western world is currently France, with extremely elevated rates also in French-speaking Switzerland, Northern Italy, Central and Eastern Europe (especially Hungary) and parts of Latin America. Rates are elevated amongst both men and women throughout South Asia. In the USA incidence rates are two-fold higher in Black men than White men (1981). Very high rates in the IARC database for Melanesia, presumably associated with areca nut and tobacco habits, are based on small numbers and need confirmation (730,1981).

The high incidence rates in Australasia are explained by lip cancer in fair-skinned races which has a comparatively low mortality rate.

Much of Europe and Japan is experiencing alarming rises in incidence, with a strong cohort effect, those born from approximately 1930 onwards showing significantly increased incidence and mortality. In North America there are statistically significant falls in Whites, but Blacks continue to show worse outcomes. Globally, with the exception of the most highly specialized treatment centres, survival rates have not improved for decades.

Significant increases in incidence in younger subjects, particularly males, have been reported from many western countries in recent decades (1534,2259).

Etiology
Tobacco smoking and alcohol
The dominant risk factors are tobacco use and alcohol abuse, which are strongly synergistic (228). Alcohol and tobacco account for 75% of the disease burden of oral and oropharyngeal malignancies in Europe, the Americas and Japan (227,1862). For the highest levels of consumption compared to the lowest ones relative risks from 70 to over 100 have been shown (287,1811). Relative risks in case-control studies showing a supermultiplicative effect in the oral cavity, between additive and multiplicative in the oesophagus, and multiplicative in the larynx, reflecting degree of contact with both these agents at these sites (797).

Most of the rise in western countries in recent years has been attributed to rising alcohol consumption in northern Europe (1597 and rises in tobacco consumption in parts of southern Europe. Significant risk increases have also been reported amongst non-drinking smokers and, to a lesser extent, non-smoking heavy
might operate, in part, on the same criti-
carinogenesis at these sites.

Prevention Recent work on risk factors in younger
cases emphasises the importance of early and heavy tobacco and alcohol
use, the protective effect of diets rich in fresh fruits and vegetables, but with a
substantial minority without these established risk factors [1534].
The protective effect of diets rich in trace elements and antioxidant vitamins is well
demonstrated in many countries, especially in Italian studies [1628,2563].
Though more controversial, a contribution
from poor oral hygiene is also suggested [108,2548].

Second primary tumours It has been recognised for a long time that
patients with oral cancer are at risk of second tumours in the upper aerodiges-
tive tract. This has been reported to occur in 10-35% of cases [2676]. These may be
synchronous with the index tumour or, if occurring after an interval of longer than
six months are described as metachro-
nous. Recurrence of the index tumour after treatment can be diagnosed by the
pathologist where the tumour is in deeper tissue and not associated with the epithelial
surface. However, the most frequent situation of second tumours is when they
arise from surface epithelium adjacent to the treated index tumour. On morphologi-
cal grounds these are diagnosed as sec-
ond primary tumours. The increasing use
of molecular biological techniques has
allowed distinction to be made between molecularly distinct second primary
tumours and second field tumours
derived from the same genetically altered
field as the index tumour [248].

Localization Tumours may arise in any part of the oral
cavity. The most common sites vary geo-
graphically reflecting different risk factors.
Lip SCC arise almost exclusively on the
lower lip. Within the oral cavity, the
subsites at which tumours may be located include: buccal mucosa, upper and
lower gingiva, hard palate, anterior two-
thirds of the tongue, including dorsal, ventral and lateral surfaces, and the floor
of mouth. Many tumours are large at pres-
etation and the tumour site is then
recorded as essentially the centre of the
tumour. Analysis of small symptomless
tumours shows the highest frequency in
floor of mouth, ventrolateral tongue and
soft palate complex [1655]. This suggests
that tumours arise at these sites, but
spread preferentially to involve other sites
such as tongue, being then recorded
as lingual lesions. The clinical relevance
of this observation is to emphasise the

Human papillomavirus (HPV) infection HPVs, especially those genotypes of
known high oncogenic potential in uterine
cervix and skin such as HPV 16 and 18,
are found in a variable but small propor-
tion of oral, and up to 50% of tonsillar and
oropharyngeal SCCs, especially the tons-
il. Recent studies suggest that HPV may
be responsible for a small fraction of oral,
and up to 40% of oropharyngeal, cancers [888,1077]. This has lead to speculation
that HPV infection, perhaps arising from
oral/genital contact, might be important in
some cases [2284]. Of interest is the
observation that HPV-containing cancers
at these sites do not generally show TP53
mutations, contrary to HPV DNA-negative
cancers [660,1077]. It is well known that
HPV 16 E6 protein inactivates p53 pro-
tein, suggesting that HPV and smoking
might operate, in part, on the same criti-
importance of close examination of high-risk sites. The oropharynx consists of the base of the tongue (posterior third), vallecula, tonsil with tonsillar fossae and pillars, glossotonsillar sulci, posterior wall and superior wall composed of the inferior surface of the soft palate and the uvula. The most common oropharyngeal site of involvement for SCC is the base of tongue.

**Clinical features**

**Signs and symptoms**

Patients with small oral and oropharyngeal SCC are often asymptomatic or may present with vague symptoms and minimal physical findings. Hence, a high index of clinical suspicion is needed to diagnose small lesions, especially if the patients have tobacco and alcohol habits. Patients may present with red lesions, mixed red and white lesions, or white plaques. Co-existing white plaques (leukoplakia) may be observed adjacent to carcinomas and this implies an origin in a pre-existing white lesion though the prevalence of this association varies considerably in different populations. However, most patients present with signs and symptoms of locally advanced disease. The clinical features may vary according to the affected intraoral subsite. Mucosal growth and ulceration, pain, referred pain to the ear, malodour from the mouth, difficulty with speaking, opening the mouth, chewing, difficulty and pain with swallowing, bleeding, weight loss, and neck swelling are the common presenting symptoms of locally advanced oral and oropharyngeal cancers. Occasionally, patients present with enlarged neck nodes without any symptoms from oral or oropharyngeal lesions. Extremely advanced cancers present as ulceroproliferative growths with areas of necrosis and extension to surrounding structures, such as bone, muscle and skin. In the terminal stages, patients may present with orocutaneous fistula, intractable bleeding, severe anaemia and cachexia.

Cancer of the buccal mucosa may present as an ulcer with indurated raised margin, exophytic or verrucous growth or with the site of origin depending upon the preferential side of chewing and placement of betel quid. In advanced stages, these lesions infiltrate into the adjacent bone and overlying skin. Cancer of the tongue may appear as a red area interspersed with nodules or as an ulcer infiltrating deeply, leading to reduced mobility of the tongue. These tumours are
Squamous cell carcinoma (SCC) may arise as a red area, a small ulcer or as a papillary lesion. Most patients present with discomfort or irritation at the site of the tumour. Advanced stages are associated with drooling. Cancers of the lower lip usually arise in the vermillion border and appear as a crusty indurated or ulcerated lesion. Cancers of the upper lip are rare, often originate on the skin and spread to the mucosa. Cancer of the gingiva usually presents as an ulceroproliferative growth. Tumours of the alveolar ridge may occasionally present as difficulty in wearing denture plates or as loosening of teeth associated with pain and bleeding during brushing of teeth. Tumours of the hard palate often present as papillary or exophytic growths, rather than a flat or ulcerated lesion. Cancer of the soft palate and uvula often appear as an ulcerative lesion with raised margins or as fungating masses. Tonsillar cancers generally appear as an exophytic or ulcerative lesion. Sometimes they can present as enlarged neck nodes without any other signs and symptoms. Cancer of the base of tongue presents late in the course of the disease as a grossly ulcerated, painful, indurated growth. More than two-thirds of the patients with buccal mucosal and gingival cancers in South Asia present with submandibular lymph node enlargement. More than three fourths of patients with tongue, floor of mouth and oropharyngeal cancers in South Asia present with neck swellings implying clinically obvious lymph node metastasis. In the West lymph node involvement is common at presentation in oropharyngeal SCC.

Imaging

Intraoral and dental radiographs, in combination with orthopantomography, may help in identifying involvement of the underlying bone. Three-dimensional imaging with computed tomography (CT) and magnetic resonance imaging (MRI) is frequently used to supplement the clinical evaluation and staging of the primary tumour and regional lymph nodes. CT scan or MRI give more information about the local extent of the disease and also help to identify lymph node metastases. CT scanning is useful in evaluating involvement of cortical bone. MRI is more informative when evaluating the extent of soft tissue and neurovascular bundle involvement. The combination of soft tissue characterisation and anatomical localization afforded by CT and MRI make them valuable tools in the
preoperative assessment of patients with oral or oropharyngeal cancers. Distant metastasis from oral and oropharyngeal cancer is uncommon at presentation. At minimum, a routine radiograph of the chest is performed to rule out lung metastases.

**Relevant diagnostic procedures**

Optimal therapy and survival from oral cancer depend on adequate diagnosis and assessment of the primary tumour and its clinical extent. Physical examination should include visual inspection and palpation of all mucosal surfaces, bimanual palpation of the floor of the mouth, and clinical assessment of the neck for lymph node involvement. The diagnosis is confirmed by biopsy. The specimen is taken from the clinically most suspicious area, avoiding necrotic or grossly ulcerated areas, and more than one biopsy site may need to be chosen. In patients with enlarged cervical lymph nodes and an obvious primary in the oral cavity or oropharynx, the biopsy is always taken from the primary site and not from the lymph node. In such situations, fine needle aspiration cytology may be carried out to verify the involvement of the node. If no obvious primary site is found in patients presenting with neck nodes, fine-needle aspiration of the lymph node can be performed to help establish the diagnosis. In patients for whom fine needle aspiration is non-diagnostic and SCC is strongly suspected, excisional lymph node biopsy is a last resort, as subsequent curative therapy may be compromised by this procedure. The search for an occult primary tumour may include direct pharyngolaryngoscopy with biopsy of high-risk sites like base of tongue, nasopharynx, and usually a diagnostic tonsillectomy, as well as other imaging modalities. Open lymph node biopsy is carried out only when the lesion cannot be identified by aspiration biopsy or in patients with suspected lymphoma. Patients with SCC of the oral cavity or oropharynx have a risk of multiple primary tumours in the pharynx or larynx, as well as in the tracheobronchial region and oesophagus so routine panendoscopy is often performed to evaluate these sites.

**Tumour spread and staging**

Staging is carried out according to the TNM classification [947,2418]. Recent additions to the coding have been provided for micrometastases, isolated tumour cells, findings in sentinel nodes and tumour detection by molecular methods. Some of these are discussed in the following sections. Local spread of oral SCC, in the early stages, is relatively predictable in tissues that have not been previously irradiated. It is influenced by local anatomical features. Lip SCC spreads superficially and then into deeper tissues. Floor of mouth SCC spreads superficially rather than in depth, being unlikely to invade into the mylohyoid muscle or the sublingual gland until a late stage. Tumour involving the lateral margin of tongue, whether arising there directly or by superficial spread from the floor of mouth, tends to spread in depth. The intrinsic muscles of tongue run in small bundles in all directions such that invading tumour encounters some muscle running at right angles to the surface. The line of least resistance to tumour spread is therefore along these muscle bundles and into the tongue. Tumours of palate spread superficially rather than in depth and this is also true for more posterior tumours of the oropharynx. For most oral SCC other than tongue, the extent of spread in an area can be predicted from the extent of surface involvement. Tongue and tonsil tumours can spread beneath intact normal appearing surface, giving a larger area of tumour involvement. Spread of oral SCC into bone is a frequent problem. The mandible is involved much more frequently than the
maxilla. In dentate jaws the usual route of entry into mandible is along the periodontal ligament. In edentulous areas of mandible the tumour spread is through the crest of the alveolus directly into the marrow spaces between trabeculae of cancellous bone [1682]. This occurs because of failure of formation of an intact cortex of alveolar bone as resorption of edentulous alveolus progresses. Tumours in the mandible can involve the inferior alveolar nerve [1683] with a particular likelihood of spread posteriorly along the nerve, sometimes extending well beyond the mandibular foramen. Cancers arising in gingiva or alveolus and those involving these sites by extension from adjacent sites are unlikely to invade into the mandible other than by periodontal liga-

Spread in previously irradiated tissues
Tumour spread in previously irradiated soft tissues tends to be more extensive and less predictable than in normal tissues and as a consequence requires more extensive surgery if excision is attempted. Tumour invasion into irradiated mandible tends to occur wherever the tumour approaches bone, often at multi-

Lymphatic spread
Spread to local lymph nodes worsens the prognosis in oral and oropharyngeal cancer. The mechanism of spread from the primary site to lymph nodes is almost always by embolism. Permeation in lymphatics adjacent to tumours is uncommon and it is debatable if this spread extends as far as lymph nodes. Once tumour is present in the neck, however, spread between nodes may be embolic or by permeation. The lymph nodes in the neck are divided into levels. The lymphatic drainage from different head and neck sites is relatively predictable [1789]. Levels at high risk for metastasis from oral cavity SCC are Levels I, II and III, and to a less-

Squamous cell carcinoma 173

els of involvement in many patients [2817]. Bilateral spread to the neck is likely to occur from tumours involving the midline, especially tumours of posterior tongue or soft palate. Extracapsular spread of tumour involving lymph nodes is associated with a poor prognosis [2819].

There have been many studies attempting to predict the presence of lymphatic spread from features of the primary tumour (872,2820). Tumour size and site are relevant. Tumour differentiation is not a reliable predictor. The pattern of the invasive front is a useful predictor in that a non-cohesive front is associated with increased likelihood of metastasis. Other factors associated with increased risk of metastasis are perineural spread at the invasive front, lymphovascular invasion and tumour thickness. The tumour thick-

Haematogenous spread
Until relatively recently, haematogenous spread of oral and oropharyngeal cancer has been regarded as less important than local and lymphatic spread. However, its importance is increasing as loco-regional control improves. Blood borne spread most often involves lung [754,1958]. The best predictor of the likelihood of this spread is involvement of the neck at multiple levels. This suggests that the route of entry of tumours into the circulation is most often via the large veins in the neck and that haematogenous spread is in effect tertiary spread following extracapsular spread from neck nodes.

Sentinel node biopsy
This is currently an experimental technique [2057] that is under active evaluation by prospective clinical trials and it is not practised at all centres. It is a techni-

cue used primarily for staging a clinical-

ly N0 neck, in an effort to avoid a neck dissection. If a clinically N0 neck is followed untreated until tumour development occurs, the prognosis can be very poor [57,977]. Studies on the incidence of occult metastases in N0 necks [753] have shown tumour spread in only a small minority of patients. Therefore, if neck dissection is undertaken either prophylacti-

cally or as a staging procedure, on patients with N0 necks, a large majority will have unnecessary surgery, as the neck will be found to be free from tumour. The sentinel node is the first draining lymph node from a tumour. It is assumed that if the sentinel node can be shown to be free from tumour, then the lymphatic basin is free from tumour and neck dissection is not required. By contrast, sen-

tinel node positive patients can be select-

ed for further therapy. Sentinel nodes are identified by a combination of lymphoscintigraphy and injection of blue dye in the tumour bed and then sampling draining nodes identified. In reality, more than one sentinel node is found in many cases [2345] indicating that tumours drain to more than a single first echelon node, presumably from different parts of the tumour. Sampled sentinel nodes should be fully examined by the pathologist. This usually involves bisecting the node in the largest diameter and then undertaking extensive sampling. Some pathologists undertake frozen sections on bisected fresh nodes. If this is done it is important to use a tech-

nique whereby the cut surface is frozen on a flat surface and only early sections are examined. This is to ensure that as little node as possible is examined at this stage in order not to compromise full examination of the node. Paraffin processed blocks are then examined with H and E sections of the early sections of the blocks. If these show no tumour, more detailed sampling with immunocytochem-

istry for cytokeratins and sampling through the block is required. True serial sectioning is impracticable for routine use. A compromise is step sectioning at intervals of 150µm with examination of H and E sections and AE1/3 reacted sec-

tions [2202]. The importance of these sections is that suspicious areas on immunocytochemistry can be identified in the H and E sections. These may be viable tumour cells, but other possible causes of cytokeratin positivity, such as inclusion of normal salivary gland epithelium or thyroid follicles, either occult metastases or lateral aberrant thyroid, need to be identified. Another not infre-

quent finding is areas of cytokeratin posi-

tivity which on H and E appear as dense-

ly eosinophilic apparently non-viable tumour cells.
Interpretation of sentinel nodes can demand considerable pathological expertise. The outcome of the pathological assessment may be the presence of metastasis; micrometastasis, less than 2mm diameter tumour deposits, or isolated tumour cells (2477). Micrometastasis has been defined (1073) as cells which have arrested and implanted. These may be in contact with a vessel or lymph sinus wall or may be extravascular. Single or small clusters of cells within lymph or blood vessels, but not in contact with the wall are defined as isolated tumour cells.

**Histopathology**

The histological features of SCC have been discussed in Chapter 3 on tumours of the hypopharynx, larynx and trachea. The findings in the oral cavity and oropharynx do not differ significantly from those of the larynx and hypopharynx. A minority of oral and oropharyngeal cancers show different histological subtypes that can be associated with differences in prognosis. These are discussed below. It is clearly important that pseudo epitheliomatous hyperplasia (PEH) is distinguished from SCC. PEH can occur in mucosa overlying a granular cell tumour, in necrotising sialometaplasia and in papillary hyperplasia of palate. PEH occurring with mucositis, particularly after irradiation, may be difficult to distinguish from squamous cell carcinoma.

The majority of cases of SCC present no difficulty in diagnosis for the experienced pathologist. However, the recognition of the earliest stages of invasion can be problematic. No consistent guidelines for this exist. The deepest layers of the epithelium and the interface between the epithelium and the lamina propria need to be examined in detail. This is frequently made more difficult where there is a prominent inflammatory infiltration.

Relevant features include the loss of a histologically well-defined interface, described previously as loss of basement membrane and disturbed architecture of the basal layers of the epithelium, particularly the replacement of basal cells by larger irregular cells with cytoplasmic processes extending into connective tissue. In some cases the degree of cytological atypia and mitotic feature may suggest malignancy, but these are not always present. To an extent the judgement about early invasion is subjective and it can be important for the pathologist to communicate the difficulty in interpretation to the clinician. Some pathologists will indicate that while no unequivocal evidence of invasion is demonstrated, they nevertheless feel that the lesion should be regarded as early invasive carcinoma.

**Somatic genetics**

There is some variation in the genetic profile of oral and oropharyngeal SCC that reflects the site-specific impact of various casual agents and differences in clinical presentation. The carcinogens in tobacco smoke, for example, increase the prevalence and spectrum of TP53 mutations (268). Compared to carcinomas that arise in patients who smoke, carcinomas in patients who have never smoked harbour fewer p53 mutations, disproportionately involve women, typically arise from the oral tongue, and affect very young or very old patients (1351,2258). For carcinomas of the oropharynx, oncogenic human papillomavirus (HPV), particularly the HPV-16 subtype, is an important causative agent: More than 50% of oropharyngeal carcinomas harbour integrated HPV DNA (60,888,1999). The E6 and E7 viral oncoproteins bind and inactivate the TP53 and retinoblastoma gene products respectively, disengaging two of the more critical pathways involved in cell cycle regulation (2788).

These HPV-positive oropharyngeal tumours compose a distinct pathological entity with its own clinical spectrum and basaloid morphology (888,1012,2072), illustrating the emerging role of genetic characterization as a potential means of determining prognosis and influencing management (1691).

Genetic evidence has clarified the vague concept of “field cancerization”. Most, if not all, multiple primary carcinomas of the upper aerodigestive tract derive from a common clonal progenitor cell that undergoes a common early genetic alterations (187,2271). Genetic evidence has helped account for the perplexing problem of local tumour recurrence following seemingly complete tumour resection. In many instances, local tumour recurrence reflects extension of genetically damaged cells beyond the clinical and microscopic boundaries of carcinoma to the margins of surgical resection (268,1626, 1983,2777).

Microsatellite analysis of exfoliated cells swabbed or rinsed from the oral cavity of patients with head and neck squamous carcinomas consistently harbour genetic changes that are identical to those in the primary tumours, suggesting a non-invasive test for specific DNA-sequence variants in saliva as a means of identifying patients with pre-invasive or invasive neoplasms (2430). Clonal genetic changes identical to those found in primary head and neck SCC have been identified in circulating plasma or serum, suggesting a mechanism for early cancer detection and tumour surveillance (1853). The use of highly sensitive genetic assays for detecting rare cancer cells at the margin of tumour resection shows promise for predicting the likelihood of tumour recurrence (268,1983).

**Prognosis and predictive factors**

Tumour size and nodal status are the most significant prognostic factors (2060). Histological grade correlates poorly with patient outcome (1292,2195). The value of grading improves when only the deeply invasive margins of the tumour are evaluated (291,292,1927, 2818). Tumours invading with pushing borders are less aggressive than tumours showing a noncohesive front showing diffuse spread with tiny strands or single cells. (1325,2132,2342,2653, 2841) Major risk factors that adversely influence prognosis...
are two or more positive regional nodes, extracapsular extension of nodal disease, or positive margins of resection (1429). Other important histologic features associated with poor prognosis are tumour thickness and vascular invasion. Molecular markers with unequivocal prognostic and/or predictive significance have not been identified (428,1052,1561,2106).

Verrucous carcinoma

ICD-O code 8051/3

Although uncommon, 75% of all cases of VC occur in the oral cavity. It is an exophytic, warty, slowly growing variant of SCC with pushing margins. It typically involves older males (950,1251,1677,1695,2621). Chronic smokeless tobacco use is accepted as the primary etiological factor for oral VC. Human papillomavirus subtypes 16 and 18 have been identified in up to 40% of oral VC (1927,2349). Oral VC begins as a well-demarcated, thin white keratotic plaque which quickly thickens and develops papillary (blunted tips) or verruciform (pointed tips) surface projections. Occasional lesions present as erythematous or pink papular masses. The colour depends on the amount of keratin produced and the degree of host inflammatory response to the tumour. This cancer almost always remains broad-based or sessile and can become quite extensive from lateral growth by the time of diagnosis. Rare fungating examples, however, may appear to be somewhat pedunculated. Smokeless tobacco keratosis (tobacco pouch) is often seen on adjacent mucosal surfaces in patients who chew tobacco or use snuff. Unless the tumour is infected or is encroaching on alveolar nerves in the jawbones, VC is an asymptomatic lesion. Surface ulceration and haemorrhage are not seen, unless a focus of SCC is present in the mass. VC consists of thickened club-shaped papillae and blunt stromal invaginations of well-differentiated squamous epithelium with marked keratinization. The squamous epithelium lacks the usual cytologic criteria of malignancy, and by morphometry, the cells are larger than those seen in SCC (489). Mitoses are rare, and observed in the basal layers; DNA synthesis (S-phase) is also limited primarily to the basal layers (737). VC invades the stroma with a pushing, rather than infiltrating border. Dense lymphoplasmacytic host response is common. Intraepithelial microabscesses are seen, and the abundant keratin may evoke a foreign body reaction. The surrounding mucosa shows progressive transition from hyperplasia to VC. A downward dipping of epithelium often “cups” the VC periphery, and is the ideal site for deep biopsy (174,1192). With extensive surgical removal, and without neck dissection, the 5-year disease-free survival rate is 80-90%, although 8% of patients require at least one additional surgical procedure during that time (1870,1927). Treatment failures usually occur in patients with the most extensive involvement or in those unable to tolerate extensive surgery because of unrelated systemic diseases. No molecular or other markers have yet shown prognostic significance for oral VC. However, one-fifth of these tumours contain a co-existing SCC which may not be identified without extensive histologic sectioning (1927). Such hybrid tumours have a greater tendency to recur locally and a slight tendency to metastasize to the ipsilateral neck.

Basaloid squamous cell carcinoma

ICD-O code 8083/3

This is uncommon in the oral cavity, slightly more common in the oropharynx. It is described in the chapter on tumours of the hypopharynx, larynx and trachea.

Papillary squamous cell carcinoma

ICD-O code 8052/3

This is rarely recognized in the oral cavity and oropharynx other than as a component of a large SCC. It is described in the chapter on tumours of the hypopharynx, larynx and trachea.

Spindle cell carcinoma

ICD-O code 8074/3

This unusual variant is more common in the larynx than in the oral cavity and oropharynx, and is described in detail in the chapter on tumours of the hypopharynx, larynx and trachea.
Lymphoepithelial carcinoma

Definition
Lymphoepithelial carcinoma (LEC) is a poorly differentiated squamous cell carcinoma (SCC) or undifferentiated carcinoma, accompanied by a prominent reactive lymphoplasmacytic infiltrate. The morphological features are indistinguishable from those examples of nasopharyngeal nonkeratinizing carcinoma with a rich lymphoplasmacytic infiltrate.

ICD-O code
8082/3

Epidemiology
LEC is rare at these sites, and accounts for 0.8-2% of all oral or oropharyngeal cancers (1339,2741). See Chapter 2.

Etiology
Epstein-Barr virus (EBV) has been tested in only a limited number of cases (819,856,1802,1875,2405), but it appears that tumours occurring in Chinese are usually positive for EBV, while those occurring in Caucasians are usually negative. The racial difference in the association with EBV is similar to LEC occurring in the major salivary glands (see Chapter 5).

Clinical features
The patients present with an intra-oral mass, which may be ulcerated. Some tumours can be bilateral (801,2038). A proportion of patients present with neck mass due to regional lymph node involvement (119).

Location and metastatic spread
More than 90% of all oral and oropharyngeal LEC occur in the tonsil and tongue base areas. The remaining cases are found in the palate and buccal mucosa (444,694,2822). The tumour has a high propensity for regional cervical lymph node involvement (approximately 70% of cases at presentation) (119,444,1339). Distant metastasis tends to occur in the liver and lung (119).

Histopathology
LEC of the oral cavity and oropharynx shows morphologic features indistinguishable from its nasopharyngeal and sinonasal counterparts. The surface epithelium is often intact. The tumour is invasive, and comprises syncytial sheets and clusters of carcinoma cells with vesicular nuclei, prominent nucleoli and ill-defined cell borders. A rich lymphoplasmacytic infiltrate is present within the tumour islands and the surrounding stroma, which may appear desmoplastic. The tumour cells are immunoreactive for pan-cytokeratin and epithelial membrane antigen. EBV encoded RNA (EBER) has been demonstrated by in-situ hybridization in oral / oropharyngeal LEC occurring in Chinese patients.

Prognosis and predictive factors
LEC of the oral cavity and oropharynx are radiosensitive, and in a high percentage of cases local control can be achieved even in the presence of regional lymph node metastasis (1339). Local, regional and distant failures occur in 3%, 5% and 19% of cases respectively (444). Distant metastasis is associated with a poor prognosis.

Fig. 4.13 A Lymphoepithelial carcinoma of the tonsil. The tumour infiltrates beneath an intact surface epithelium. In this example, the tumour islands are obscured by the heavy lymphoplasmacytic infiltrate. B Sheets and islands of tumour cells intimately admixed with lymphocytes and plasma cells. C Lymphoepithelial carcinoma of the palate. Carcinoma cells exhibit indistinct cell borders, pale chromatin and distinct nucleoli. Many lymphocytes are found among the carcinoma cells.
Epithelial precursor lesions

The pathologic assessment of precursor lesions is similar throughout the upper aerodigestive tract. It is described in detail in the Chapter 3 on tumours of the hypopharynx, larynx and trachea (page 140).

**Clinical features**
The principal oral and oropharyngeal lesions which may be precursor lesions are white patches (leukoplakia) and red patches (erythroplasia/erythroplakia) or mixed red and white lesions. The majority of leukoplakias will not show dysplasia and correspond to the hyperplasia category. Red and mixed lesions (speckled leukoplakia) show a higher frequency of dysplasia, often of higher grade. The majority of leukoplakias will not undergo malignant change and may even regress particularly if apparent aetiologic factors are removed.

**Histopathology**
The epithelium of precursor lesions may be thick, but in the oral cavity it can also be atrophic. By definition, there is no evidence of invasion. The magnitude of surface keratinisation is of no importance. Allocation to categories within each of the classifications requires consideration firstly of architectural features and then of cytology.

**Hyperplasia**
Hyperplasia describes increased cell numbers. This may be in the spinous layer (acanthosis) and/or in the basal/parabasal cell layers (progenitor compartment), termed basal cell hyperplasia. The architecture shows regular stratification without cellular atypia.

**Dysplasia, / squamous intraepithelial neoplasia / atypical hyperplasia**
When architectural disturbance is accompanied by cytologic atypia, the term dysplasia applies. The terms squamous intraepithelial neoplasia (SIN) and atypical epithelial hyperplasia are used synonymously. There is a challenge in the recognition of the earliest manifestations of dysplasia and no single combination of the above features allows for consistent distinction between hyperplasia and the earliest stages of dysplasia. Dysplasia is a spectrum and no criteria exist to precisely divide this spectrum into mild, moderate and severe categories.

**Mild dysplasia**
In general architectural disturbance limited to the lower third of the epithelium accompanied by cytological atypia define the minimum criteria of dysplasia.

**Moderate dysplasia**
Architectural disturbance extending into the middle third of the epithelium is the initial criterion for recognizing this category. However, consideration of the degree of cytologic atypia may require upgrading.

**Severe dysplasia**
Recognition of severe dysplasia starts with greater than two thirds of the epithelium showing architectural disturbance with associated cytologic atypia. However, as noted in the previous para-

### Table 4.01 Classification schemas that histologically categorize precursor and related lesions

<table>
<thead>
<tr>
<th>2005 WHO Classification</th>
<th>Squamous Intraepithelial Neoplasia (SIN)</th>
<th>Ljubljana Classification Squamous Intraepithelial Lesions (SIL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Squamous cell hyperplasia</td>
<td>SIN 1</td>
<td>Basal/parabasal cell hyperplasia*</td>
</tr>
<tr>
<td>Mild dysplasia</td>
<td>SIN 2</td>
<td>Atypical hyperplasia**</td>
</tr>
<tr>
<td>Moderate dysplasia</td>
<td>SIN 3***</td>
<td>Atypical hyperplasia**</td>
</tr>
<tr>
<td>Severe dysplasia</td>
<td>SIN 3***</td>
<td>Carcinoma in-situ</td>
</tr>
<tr>
<td>Carcinoma in-situ</td>
<td>SIN 3***</td>
<td>Carcinoma in-situ</td>
</tr>
</tbody>
</table>

* Basal/parabasal cell hyperplasia may histologically resemble mild dysplasia, but the former is conceptually benign lesion and the latter the lower grade of precursor lesions.
** 'Risky epithelium'. The analogy to moderate and severe dysplasia is approximate.
*** The advocates of SIN combine severe dysplasia and carcinoma in-situ.

### Table 4.02 Criteria used for diagnosing dysplasia

<table>
<thead>
<tr>
<th>Architecture</th>
<th>Cytology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Irregular epithelial stratification</td>
<td>Abnormal variation in nuclear size (anisonucleosis)</td>
</tr>
<tr>
<td>Loss of polarity of basal cells</td>
<td>Abnormal variation in nuclear shape (nuclear pleomorphism)</td>
</tr>
<tr>
<td>Drop-shaped rete ridges</td>
<td>Abnormal variation in cell size (anisocytosis)</td>
</tr>
<tr>
<td>Increased number of mitotic figures</td>
<td>Abnormal variation in cell shape (cellular pleomorphism)</td>
</tr>
<tr>
<td>Abnormally superficial mitoses</td>
<td>Increased nuclear-cytoplasmic ratio</td>
</tr>
<tr>
<td>Premature keratinization in single cells (dykeratosis)</td>
<td>Increased nuclear size</td>
</tr>
<tr>
<td>Keratin pearls within rete pegs</td>
<td>Atypical mitotic figures</td>
</tr>
<tr>
<td>Increased number and size of nucleoli</td>
<td></td>
</tr>
</tbody>
</table>
graph architectural disturbance extending into the middle third of the epithelium with sufficient cytologic atypia is upgraded from moderate to severe dysplasia.

Carcinoma in-situ
The theoretical concept of carcinoma in-situ is that malignant transformation has occurred but invasion is not present. It is not possible to recognize this morphologically. The following is recommended for the diagnosis of carcinoma in-situ: full thickness or almost full thickness architectural abnormalities in the viable cellular layers accompanied by pronounced cytologic atypia. Atypical mitotic figures and abnormal superficial mitoses are commonly seen in carcinoma in-situ.

Differential diagnosis
Reactive, regenerative or reparative squamous epithelium, for example in response to trauma, inflammation, irradiation or ulceration, may manifest atypical cytology or architectural disturbance. Nutritional deficiencies such as iron, folate, and vitamin B12, can also simulate dysplasia. Such lesions are not considered precursor lesions and should be distinguished from them. Clinical history is helpful and morphological changes suggestive of the inciting event, such as ulceration, inflammation, haemorrhage, radiation-induced mesenchymal and/or endothelial nuclear enlargement and hyperchromatism, may be present. The epithelial changes in these cases are generally less pronounced than in dysplasia.

Relevance of dysplasia.
It is reasonable to assume that the changes described in dysplasia are due to genetic changes in the epithelium occur, but it is unlikely that the mutations involved are the same ones as are associated with development of malignancy. More severe dysplasia has been traditionally believed to be associated with a greater likelihood of progression to malignancy. This might indicate that the greater the accumulation of mutations in tissue, the greater the chance that the critical mutations for malignancy will be present. The corollary is also true in that malignancy can arise from non-dysplastic epithelium [2493] presumably because these critical mutations can be present in the absence of the mutations causing dysplasia.

Genetics
There are no individual markers that reliably predict malignant transformation. The molecular biology techniques which show most promise as predictors of development of SCC are large scale genomic status (DNA ploidy) and loss of heterozygosity (LOH) at defined loci [2286]. Dysplasia has been reported to be present in from 10-25% of leukoplakias.

Table 4.03 Malignant transformation of oral leukoplakia (Reibel (2145))

<table>
<thead>
<tr>
<th>Authors/Year</th>
<th>Country</th>
<th>Material (no. of cases)</th>
<th>Observation period (years)</th>
<th>Cases with malignant transformation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pindborg et al., 1968 (2049)</td>
<td>Denmark</td>
<td>248</td>
<td>3.9</td>
<td>4.4</td>
</tr>
<tr>
<td>Silverman and Rosen, 1968 (2362)</td>
<td>USA</td>
<td>117</td>
<td>1-11</td>
<td>6.0</td>
</tr>
<tr>
<td>Kramer et al., 1970 (1368)</td>
<td>UK</td>
<td>187</td>
<td>-</td>
<td>4.8</td>
</tr>
<tr>
<td>Mehta et al., 1972 (1699)</td>
<td>India</td>
<td>117</td>
<td>10</td>
<td>0.9</td>
</tr>
<tr>
<td>Silverman et al., 1976 (2358)</td>
<td>India</td>
<td>4762</td>
<td>2</td>
<td>0.13</td>
</tr>
<tr>
<td>Bánóczy, 1977 (118)</td>
<td>Hungary</td>
<td>670</td>
<td>9.8</td>
<td>6.0</td>
</tr>
<tr>
<td>Silverman et al., 1984 (2361)</td>
<td>USA</td>
<td>257</td>
<td>7.2</td>
<td>17.5</td>
</tr>
<tr>
<td>Lind, 1987 (1517)</td>
<td>Norway</td>
<td>157</td>
<td>9.3</td>
<td>8.9</td>
</tr>
<tr>
<td>Schepman et al., 1998 (2261)</td>
<td>Netherlands</td>
<td>166</td>
<td>2.5</td>
<td>12.0</td>
</tr>
</tbody>
</table>
Ploidy studies of dysplastic leukoplakias showed that the great majority of aneuploid lesions developed SCC in the follow-up period, by contrast with 60% of tetraploid lesions and only about 3% of diploid lesions [2490]. No correlation was found between the degree of dysplasia and DNA ploidy. Similar studies on erythroplasias [2491] confirmed the high predictive potential of aneuploidy in identifying cases which progressed to SCC. Non-dysplastic white patches have also been studied [11] and although there was a much lower incidence of malignant transformation, 80% of such cases were aneuploid.

LOH studies have been undertaken contrasting oral lesions which progressed to SCC or carcinoma in-situ during follow-up with corresponding lesions which did not progress. LOH on two chromosome arms, 3p and 9p seemed to be particularly important in predicting progression [2201].

Fig. 4.15 A Moderate dysplasia. Drop shaped rete ridges, dysplasia extending to mid-third and moderate cytological changes. B Severe dysplasia into upper third of epithelium with marked cytological change. C Severe dysplasia into upper third of epithelium with prominent cytological change including abnormal mitoses. D Carcinoma in-situ. Abnormal cells seen throughout the full thickness of epithelium.
Proliferative verrucous leukoplakia and precancerous conditions

Definition

Proliferative verrucous leukoplakia (PVL) is a rare but distinctive high-risk clinical form of oral precursor lesions. Because of the lack of specific histologic criteria, the diagnosis is based on combined clinical and histopathologic evidence of progression. Sequential biopsies show progressive dysplasia and the acquisition of aberrant TP53 protein.

Clinical features

PVL is an aggressive form of oral leukoplakia with considerable morbidity and strong predilection to malignant transformation {174,1005,1797,2360}. The etiology of this entity is unknown. The condition develops initially as focal clinical hyperkeratosis (leukoplakia) that progressively becomes a wide multifocal disease with gross exophytic features {174}. The average age at diagnosis is 62 years; women are more commonly afflicted (ratio, 4:1). Typically, multiple oral sites are affected. The most common site in women is the buccal mucosa and the tongue in men. Carcinoma develops after a protracted period of time. The most common sites of the carcinoma are gingiva and tongue. PVL is characterized by high recurrence rate and histological progression. Many cases are resistant to all forms of treatment, including laser microsurgery, surgical excision and radio-and chemotherapy. Conservative management of these lesions has been unsuccessful and wide surgical excision is the best hope for control.

Other precancerous conditions

Precancerous conditions (PCs) are generalized clinical states associated with a significantly increased risk for SCC. Epithelial atrophy, increased mitotic activity and impaired epithelial repair mechanisms are fundamental to PCs of different etiology.

Iron deficiency

Originally described in the context of sideropenic dysphagia, it is an important cause of epithelial atrophy. The association of iron deficiency with oropharyngeal squamous cell carcinomas has been observed since the mid-thirties of the 20th century {21}. However, a significant decrease of cases with hypopharyngeal cancers and iron deficiency was noted in Sweden in the seventies {1433}. Few cases of oral cancer and iron deficiency have been published in the last 20 years.

Fig. 4.16 Proliferative verrucous leukoplakia (PVL) A Extensive, thick, white plaques. B Hyperplasia and dense hyperkeratosis of early PVL. C Histology from a clinical case of PVL showing verrucous surface with hyperkeratosis, hypergranulosis and a dense inflammatory infiltrate in the corium. D Same case as shown on fig. C two years later showing more florid verrucous hyperplasia illustrating the progressive nature of the condition.
Proliferative verrucous leukoplakia

Oral lichen planus

OLP is a chronic mucocutaneous immune inflammatory condition. Malignant transformation is still controversial [639, 2359]; one review reporting malignant transformation rates between 0% and 5.6% [2116]. The controversy is due to lack of uniform clinical and histological criteria for OLP and oral lichenoid lesions (OLL). The latter have also been termed interface mucositis or lichenoid mucositis. Oral lichenoid lesions have been considered by some to represent the lesion at risk if associated with dysplasia. In a recent study [2664] it was shown that all cases of malignant transformation (1.7%) involved cases of OLL and not OLP. Similarly, a study [2896] investigating whether OLP without dysplasia is premalignant by using microsatellite analysis for loss of heterozygosity (chromosomes 3p, 9p, 17p) did not support OLP as a lesion at risk.

However, until distinct clinical and histological criteria have been developed on how to differentiate OLP from OLL, both lesions have to be considered as ‘at risk for malignant transformation’.

Oral submucous fibrosis (OSF)

This chronic, progressive condition of the oral mucosa [2115] is etiologically strongly associated with the chewing of areca nut which has recently been categorized by IARC as a human carcinogen [1]. It is almost exclusively seen in ethnic groups using areca nut alone or as a component of betel quid.

Clinically there is mucosal rigidity of varying intensity due to fibroelastic transformation of the juxtaepithelial connective tissue. Fibrous bands and mucosal pallor are characteristic [498]. Histologically, there is epithelial atrophy, keratosis and dysplasia in up to 25% of cases [498]. In a population-based prospective study, in India, SCC developed in 7% of patients with OSF over a period of 17 years [1798].

Syphilis

Late stage (tertiary) syphilis associated with leukoplakia had a high risk of malignancy, but this is now largely of historical interest [1721].

Lupus erythematosus

This is a chronic autoimmune disease of unknown etiology. Carcinomas, mainly of the lips, have been described in affected individuals [2264, 2696].

Epidermolysis bullosa dystrophicans (Hallopeau-Siemens type)

This disease of the skin and oral mucosa has an autosomal dominant pattern of inheritance. Oral leukoplakia and occasional cases of SCC have been observed in association with epidermolysis bullosa [226, 2288].

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Xeroderma pigmentosum

This is a rare neurocutaneous disease with an autosomal-recessive mode of inheritance. The syndrome is caused by deficient nucleotide excision repair mechanisms [2090]. The skin, including the lips, is affected and shows epithelial atrophy and hyperpigmentation. Patients are extremely sensitive to light and show an increased predisposition to UV-associated malignancies of the skin. Carcinomas of the tongue have also been described [1306, 1994, 2704].

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**Papillomas**

**Definition**
These form a range of localised hyperplastic exophytic and polypoid lesions of hyperplastic epithelium with a verrucous or cauliflower-like morphology. Lesions of fibroepithelial hyperplasia are not generally included. Not every papilloma can be allocated to one of the diagnostic categories described below.

**ICD-O code** 8050/0

**Epidemiology**
Papillomas are common, with a prevalence of approximately 0.1%-0.5% (94, 1342,2569).

**Etiology**
HPV infection causes some papillomas [2076] and at least types 1,2,3,4,6,7,10, 11,13,16,18,30,31,32,33,35,38,45,52, 55,57,59,69,72,73 sequences have been detected in benign oral lesions. Clinically, latent HPV is common in oral mucosa and HPV DNA sequences can be detected in over 80% of individuals. There is no absolute association between the virus type and the type of papilloma [866] though focal epithelial hyperplasia is almost exclusively associated with types 13 and 32. HPV infection of oral tissue may be transmitted horizontally, including venereally, perinatally and possibly in utero [2518].

**Histopathology**
Histological differential diagnosis for all types includes lesions of fibroepithelial hyperplasia: fibroepithelial polyps, fibrous epulis and papillary hyperplasia associated with candidal infection or dentures. These have a more prominent fibrous component and no viral change. Verruciform xanthoma is a solitary lesion with a very similar clinical and histological presentation.

Extensive multiple papillomas or diffuse papillomatous change raise the possibilities of HPV lesions in immunosuppression, acanthosis nigricans, naevoid lesions and lateris, focal dermal hypoplasia, Cowden syndrome, papillary and verrucous dysplastic lesions and papillary squamous [2488] or verrucous carcinoma. Florid oral papillomatosis is a clinical term for diffuse papillomatous change of the mucosa for which no specific cause can be identified and is not a defined clinico-pathological entity.

**Squamous cell papilloma and verruca vulgaris**

**Definition**
A benign, hyperplastic wart-like localised proliferation of the oral epithelium [2076].

**Epidemiology**
Squamous papillomas are common in children and in adults in the 3rd to 5th decades but may be found at any age. There is an almost equal sex incidence with a slight male predominance.

**Etiology**
Evidence of causative HPV infection can be found in less than half of oral squamous papillomas [866,2516,2747], and these lesions are the intraoral counterpart of verruca vulgaris. Many HPV subtypes have been detected including 2,4,6,7,10,40. The presence of HPV viral components ultrastructurally and immunocytochemically indicates active viral replication in the lesion. Virus transmission appears to be mostly horizontal or by autoinoculation. Lesions in children tend to arise at anterior oral sites and the source of infection is often verruca vulgaris on the skin, particularly on the fingers. Infectivity is low.

The remainder of squamous papillomas are of unknown etiology. HPV sequences may be detected by PCR but the significance of this is unclear.
Localisation
Any oral site may be affected but the most common are the hard and soft palate, labial mucosa, tongue and gingiva.

Clinical features
Squamous papillomas are soft, pedunculated lesions formed by a cluster of finger-like fronds or a sessile dome-shaped lesion with a nodular, papillary or verrucous surface. The surface may be white or of normal mucosal colour depending on the degree of keratinization (2076). Lesions are usually single but fairly frequently multiple, particularly in children and for verruca vulgaris. Squamous papillomas grow rapidly over a period of a few months to a maximum of about 6 mm diameter and then remain a constant size.

Histopathology
Lesions are exophytic and comprise folds of hyperplastic stratified epithelium that are usually thickly para- or orthokeratinised but may be non keratinised. Squamous papillomas associated with HPV (oral verruca vulgaris) comprise a cluster of finger-like projections from a narrow base, each with a sharp keratinised tip, supported on ramifying cores of connective tissue containing dilated capillaries. Stratification of the epithelium is well ordered. Mitoses may be frequent and there may be mild anisonucleosis consistent with hyperplasia, but no atypia. The fronds are thickly keratinised, often with a prominent keratohyaline layer of large coarse granules. Small foci of HPV-infected cells (koilocytes) can usually be found in the upper prickle cell layer. These keratinocytes have crumpled, darkly stained nuclei with perinuclear haloes but appear very similar to vacuolated keratinocytes that are common in the normal oral mucosa. Koilocytes may be more frequent in early lesions. Less frequently, viral inclusions are found. Rete processes at the base often turn inwards and are symmetrical. Small foci of lymphocytic inflammation may lie in the fronds or at the base but inflammation is usually sparse unless the lesion is subject to trauma or other irritation (4,1929). HPV may be identified by immunocytochemistry or in-situ hybridisation but this is not necessary for diagnosis (2076).

Papillomas without detectable active HPV replication show more variation. They may appear identical to verruca vulgaris but without koilocytes or prominent keratohyaline granules, or form rounded broad-based dome shaped lesions similar to condyloma. The hyperplastic epithelium may form papillary exophytic fronds or arborising rete processes. Some are flat zones of acanthotic hyperplastic epithelium with increased numbers of dermal papillae similar to plane warts of the skin.

Prognosis and predictive factors
Oral verruca vulgaris may regress spontaneously, particularly in children, but responds to simple excision or ablation by laser or cryosurgery. Recurrence is unusual provided all lesional tissue is removed and there is no malignant potential.

Condyloma acuminatum

Definition
Oral counterpart of anogenital condyloma acuminatum

Synonyms
Venereal wart; venereal condyloma

Epidemiology
Lesions are usually diagnosed between the mid 2nd and 5th decade with a peak in teenagers and young adults (2916).

Etiology
Epithelial infection by HPV, most commonly types 6, 11, 16 and 18 though others have been detected (700,1380). Transmission is usually venereal or by autoinoculation from concomitant genital lesions (1975). Histological appearance is not an accurate indicator of a genital origin.

Fig. 4.22 Condyloma acuminatum. A Several sessile, cauliflower-like swellings forming a cluster. B Typical papilloma structure in condyloma showing the more rounded architecture in comparison with verruca vulgaris. Note a verrucous area on the left; many of these lesions have features of both types of papilloma.

Localisation
Most lesions arise on the labial mucosa, tongue and palate in anterior oral sites though any area may be affected (700, 2916).

Clinical features
Condylomas are painless, rounded, dome-shaped exophytic nodules up to 15 mm in diameter, larger than squamous papillomas and verruca vulgaris. They have a broad base and a nodular or mulberry-like surface that is slightly red, pink or of normal mucosal colour. Lesions may be multiple and are then usually clustered (2076,2916).

Histopathology
Condylomas are similar to squamous papillomas but with short blunt rounded fronds of hyperplastic epithelium of even length forming a smooth or nodular, flat or rounded surface. Keratin is usually absent or sparse, occasional examples show moderate keratin and are white clinically. Between the folds, crypts or clefts lined by epithelium extend close to the broad base and may be filled with keratin debris in keratinised lesions. Clusters of koilocytes identical to those described above are much commoner than in squamous papillomas and are usually a prominent feature. Unlike squamous papilloma, rete processes are bulbous and short, of even length and do not curve inwards (700,2076).

Prognosis and predictive factors
Condyloma acuminatum often responds to simple excision or ablation by laser or cryosurgery but appears to carry a higher risk of recurrence than squamous papilloma. Unlike ano-genital condyloma, there is no documented risk of malignant transformation, regardless of the presence of high-risk HPV types.
Condyloma acuminatum in children raises the possibility of sexual abuse, but non-sexual transmission is possible [1380] and probably frequent.

**Papillomas and papillomatosis in immunodeficiency**

More florid presentations of HPV-induced lesions are found in immunosuppression, particularly in HIV infection. Lesions may be larger, multiple and coalesce to form extensive patches of affected mucosa. Occasionally the entire oral mucosa may become papillomatous and some of these presentations are not easily classified. Unusual HPV subtypes and multiple HPV subtypes are more frequent in immunosuppression. Occasional lesions in HIV infection are dysplastic and are of uncertain malignant potential.

**Focal epithelial hyperplasia**

**Definition**
Multiple oral papillomas induced by HPV 13 and 32

**Synonym**
Heck disease

**Epidemiology**
This is primarily a disease of children, adolescents and young adults. Originally described in Inuit and native Americans [69] but now recognised worldwide. The condition is endemic in some countries and prevalence may be as high as 40% of children in localised areas [94,332,1014].

**Etiology**
Infection by HPV types 13 and 32.

**Localization**
All areas of the oral cavity may be affected but the lesions are most common on the labial and buccal mucosa and the tongue [69,332,1014].

**Clinical features**
Typically there are multiple asymptomatic lesions, each a soft rounded or flat plaque-like sessile swelling with a slightly nodular surface. They are usually pink in colour or sometimes white, and 2-10mm in diameter. Lesions develop in clusters or confluent patches [332]. Individual lesions may appear and disappear during the course of the disease [1014].

**Histopathology**
The histological features are more distinctive than squamous papilloma or condyloma. Each lesion is a slightly raised or rounded sessile swelling formed by a sharply demarcated zone of epithelial acanthosis, similar to condyloma acuminatum but with a less prominent papillomatous structure. The bulk of lesion is formed by exophytic acanthosis, without formation of well-defined projections of epithelium and the lesion contains minimal connective tissue papillae. Koilocytes similar to those of squamous papilloma are usually present and, in addition, there are usually characteristic "mitosoid bodies", which are nuclei with coarse clumped heterochromatin resembling a mitotic figure. Mitosoid bodies are characteristic but not specific for focal epithelial hyperplasia. The base of the lesion is flat and level with the adjacent epithelium without rete process enlargement [332,2076]. HPV may be detected on immunocytochemistry or by in-situ hybridisation but this is not necessary for diagnosis if the clinical presentation is typical [2076].

**Genetic susceptibility**
Familial clustering and endemic areas may result from horizontal transmission.

**Prognosis and predictive factors**
The condition appears to resolve spontaneously after a period of years and is rarely found in adults. It has no malignant potential.
**Definition**
A benign tumour of soft tissues which most often arises in the tongue and is thought to be of Schwann cell origin. It is composed of a poorly demarcated accumulation of plump granular cells which are often intimately associated with skeletal muscle.

**ICD-O code** 9580/0

**Synonym**
Granular cell myoblastoma

**Epidemiology**
Granular cell tumours are rare. Approximately 50% of all lesions arise in the head and neck and over half of these are found in the tongue. They arise in all age groups, with a peak between 40 and 60 years. In about 10-20% of patients the lesions are multiple. Females are affected more often than males with an M/F ratio of 2:1.

**Etiology**
No etiological factors are known. The lesion is thought to arise from Schwann cells. The granularity may be a senescent change associated with accumulation of lysosomes.

**Localization**
Granular cell tumours may arise in the skin, soft tissues, breast and lungs, but over 50% involve the head and neck and the tongue is the most common single site. Oral lesions may also be found in the buccal mucosa, floor of oral cavity or palate. Lesions may be multiple, affecting more than one intraoral site, or involving oral and extraoral sites [477]. Rare lesions have been reported in the salivary glands [331].

**Clinical features**
The lesion typically presents as a smooth, sessile mucosal swelling 1-2 cm in diameter with a firm texture. The overlying epithelium is of normal colour or may be slightly pale. Occasionally there is candidal infestation of the superficial epithelium and the lesion may then present as a discrete, white plaque.

**Macroscopy**
Tumours are usually 1-2 cm in diameter with a smooth surface. The cut surface shows a poorly demarcated lesion which is pale yellow or cream and firm on cutting.

**Histopathology**
The lesion is composed of plump eosinophilic cells with central small dark nuclei and abundant granular cytoplasm. The cells may be polygonal or elongated and have indistinct cell membranes, often giving the impression of a syncytium. The lesion is not encapsulated and the granular cells extend into adjacent tissues, typically skeletal muscle, where they appear to merge with muscle cells [477,2791]. Granular cells extend up to the epithelium, often forming small islands in the connective tissue papillae. The granules stain positively with periodic acid Schiff (PAS). A characteristic feature of granular cell tumour is that in up to 30% of cases the overlying epithelium shows pseudoepitheliomatous hyperplasia that may be misdiagnosed as carcinoma.

**Immunoprofile**
The lesion is strongly and uniformly positive for S-100 protein. Cells also express neuron-specific enolase, calretinin, inhibin-alpha and PGP 9.5, and show fine granular cytoplasmic positivity for the lysosome related antigen CD68 [764, 2791].

**Prognosis and predictive factors**
Granular cell tumours are benign and rarely recur, even after conservative removal. Occasional lesions have behaved aggressively and malignant granular cell tumours have been described.
Fig. 4.27 Granular cell tumour. **A** The granular cells frequently extend close to the overlying epithelium, but do not fuse with it. **B** The granular cells infiltrate widely and often appear to merge with striated muscle cells. **C** The granules are PAS positive (Periodic acid Schiff stain). **D** The granular cells are strongly and uniformly positive for S-100 protein.
Keratoacanthoma

Definition
Keratoacanthoma is a benign tumour that is believed to arise from the epithelium of hair follicles.

ICD-O code 8071/1

Synonyms
Molluscum sebaceum, molluscum pseudocarcinomatous, self-healing primary squamous carcinoma, tumour-like keratosis, idiopathic cutaneous pseudo-epitheliomatous hyperplasia.

Epidemiology
Keratoacanthoma occurs more often in whites, and is almost twice as frequent in men as in women. Although they have been seen in infants, keratoacanthomas are rare in persons under 20 years of age and the peak incidence is between the sixth and seventh decade [881].

Etiology
Interestingly, the uptake of carcinogens (e.g. via particular smoking habits) may be relevant in human tumours [641]. No other risk factors are known. The concept of a common viral origin (papillomavirus-es), popular for some years, has been abandoned.

In addition to the solitary type, clinical variants with multiple keratoacanthomas have been described, sometimes with a unilateral distribution [881]. Genetic factors may be involved in these cases, for familial clustering occurs, with multiple keratoacanthomas in affected individuals.

Localization
Keratoacanthomas preferentially occur on sun exposed hairy skin [881]. Thus, they are frequent on the skin of the face, including the lips (8% of cases), and extremely rare at hairless sites. Whether or not a “true keratoacanthoma” of the oral mucosa exists or not remains controversial [1929]. However, a small number of cases of the solitary form have been reported in intraoral sites [414,973], and mucocutaneous linings may also be affected in the generalized forms (e.g. the Ferguson-Smith, Grzybowski and Witten and Zak types) [881].

Clinical features
Keratoacanthoma is characterised by rapid growth followed by slow, spontaneous involution over several months [881]. Exact figures about regression time, however, are difficult to obtain, since the common mode of treatment is excision. The mature lesion is usually bud- or dome-shaped and is brownish or slightly reddish. Over time a central keratinous crater appears at the expense of the surrounding softer tumour tissue until finally a cup- or saucer-shape lesion develops that appears ulcerated, but is, in fact, lined by tumour epithelium and often covered with horn masses. An eruptive variant can be distinguished which is multifocal and often lacks the central keratin-filled crater. Following trauma and/or infection, true ulceration may occur, especially in areas like the lips, probably due to repeated scratching or biting. In the oral cavity, the above-described phenotypes rarely occur. Instead, the putative oral lesion mimics a broad spectrum of pseudoneoplastic and neoplastic lesions [1929].

Macroscopy
The basic gross features of epidermal lesions have been already described. However, such prototypic lesions are rarely seen in the oral cavity. Instead, as in cases at the inner side of the vulva and within the anal canal, oral keratoacanthomas present as verrucous, speckled or even ulcerated lesions. Also, they may produce deep projections, which can extend through minor salivary glands and reach the surface of underlying bone.

Histopathology
Keratoacanthomas show a verrucous surface, and underneath keratinized clefts and penetrating squamous rete processes are found with deep keratin pearls. Atypia is minimal, and mitotic figures are rare or absent. Dense inflammatory infiltrates, including granulocytes typically are found in the adjacent stroma and within the deep parts of the tumour, so that the margins seem ill defined. The
hallmark of keratoacanthoma is the overall architecture, with a cup-shaped appearance and a collar-like circumference.

A major diagnostic problem arises when destructive infiltration takes place as has been reported, including some cases in young individuals. When this kind of tumour growth occurs in the elderly, it is of course extremely difficult or even impossible to distinguish the lesion from carcinoma, particularly from carcinoma cuniculatum, which also shows minimal atypia despite its destructive growth pattern. 

Histogenesis
A large body of evidence exists pertaining to the histogenesis of keratoacanthomas. In fact, it is their origin from pilosebaceous follicles which has lead some authors to deny the existence of intraoral keratoacanthomas. This standpoint may be acceptable for sites of the oral cavity where pilo-sebaceous rudiments are rarely seen (e.g. gingiva). However, there are also cases reported in areas such as the buccal mucosa, which is a preferential site for the ectopic sebaceous glands (Fordyce spots). In addition, as also suggested for skin lesions, preprogrammed progenitor cells of the most superficial (intraepidermal) parts of the pilosebaceous unit may be sufficient as a source of (intraoral) keratoacanthoma.

Prognosis and predictive factors
Epidermal keratoacanthomas are clearly benign lesions. However, for similar tumours of the external openings (oral cavity, vulva, anal canal) there are no reliable data, since these lesions are extremely rare, present diagnostic problems and therefore are usually completely excised. Recurrences after surgical excision do not occur.
**Papillary hyperplasia**

Papillary hyperplasia is an asymptomatic nodular or papillary mucosal lesion typically seen in the palate of patients who wear dentures. Most patients wear ill-fitting dentures, wear dentures continuously (2645) or have poor denture hygiene. Lesions also arise in non-denture wearers, in xerostomia or individuals with a high arched palate. Florid and extensive presentations occur in immunosuppression (937) and HIV infection (2150). There is sessile nodular papillomatous hyperplasia of epithelium and supporting underlying fibrous tissue. There is usually parakeratinisation or less frequently orthokeratinisation. Rete processes are usually rounded or sharply defined at the base of the lesion but there may be pseudoepitheliomatous hyperplasia with keratin pearls and a poorly defined deep margin. Differential diagnosis includes diffuse HPV-induced papillomatosis, periorificial plasmacytosis (937) and verruciform xanthoma. Other multinodular lesions such as focal epithelial hyperplasia, acanthosis nigricans and Cowden syndrome appear similar histologically but have distinctive clinical presentations.

**Median rhomboid glossitis**

Median rhomboid glossitis typically forms a patch of papillary atrophy near the midline of the dorsum of the tongue at the junction of the anterior two thirds and posterior third in the region of the embryological foramen caecum. It is no longer thought to be a developmental defect but the result of chronic candidal infection (719,2825). The epithelium lacks papillae, and shows psoriasiform hyperplasia and sometimes areas of pseudoepitheliomatous hyperplasia. A mild degree of atypia may be present. Fungal hyphae are present in the superficial epithelium but are usually sparse and revealed only in multiple sections. Scarring and nodularity persist after antifungal treatment. Differential diagnosis is aided by knowledge of the specific site and includes reactive fibroepithelial hyperplasia, granular cell tumour and other nodular lesions of the tongue. Occasionally the lesion can be difficult to differentiate from squamous cell carcinoma (931, 1932), particularly when hyperplasia is extensive and epithelial processes reach or penetrate the underlying muscle.
Salivary gland tumours

Epidemiology

Tumours of the oral cavity and oropharynx account for 9-23% of all salivary gland neoplasms in major series (669, 704,2301). The most common sites are the palate (44-58%), lips (15-22%) and buccal mucosa (12-15%) (669,704,2301, 2711). Variations in these series probably reflect patterns of referral in different institutions, together with geographical and ethnic differences.

Tumours of the oropharynx are relatively uncommon and form only 1.1-3.3% of all minor gland tumours (669,704,2448). Most studies show a female to male ratio in the range of 1.2:1-1.5:1 (2711).

Location

Nearly half of all oral and oropharyngeal salivary tumours are malignant, and in some sites, such as the lower lip, tongue and floor of the oral cavity, the large majority are carcinomas. It is interesting to note that while 80-90% of labial salivary gland tumours involve the upper lip, there is a 3-5x greater risk of neoplasms in the lower lip being malignant (400,669, 704, 1871,1963). Most of the principle types of salivary gland tumour have been reported in the oral cavity. In some tumours such as canalicular adenoma, duct adenomas and polymorphous low-grade adenocarcinoma, the minor glands are by far the most frequent site of involvement. Whether there are genuine cases of intraoral Warthin tumour, or whether reported examples represent oncocytic hyperplasia and metaplasia with reactive lymphocytic infiltration, is contentious (2669).

ICD-O codes

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<th>Tumour Type</th>
<th>Code</th>
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<tbody>
<tr>
<td>Acinic cell carcinoma</td>
<td>8550/3</td>
</tr>
<tr>
<td>Mucoepidermoid carcinoma</td>
<td>8430/3</td>
</tr>
<tr>
<td>Adenoid cystic carcinoma</td>
<td>8200/3</td>
</tr>
<tr>
<td>Polymorphous low-grade adenocarcinoma</td>
<td>8525/3</td>
</tr>
<tr>
<td>Epithelial-myoepithelial carcinoma</td>
<td>8562/3</td>
</tr>
<tr>
<td>Clear cell carcinoma, NOS</td>
<td>8310/3</td>
</tr>
<tr>
<td>Basal cell adenocarcinoma</td>
<td>8147/3</td>
</tr>
<tr>
<td>Cystadenocarcinoma</td>
<td>8450/3</td>
</tr>
<tr>
<td>Mucinous adenocarcinoma</td>
<td>8480/3</td>
</tr>
<tr>
<td>Oncocytic carcinoma</td>
<td>8290/3</td>
</tr>
<tr>
<td>Salivary duct carcinoma</td>
<td>8500/3</td>
</tr>
<tr>
<td>Myoepithelial carcinoma</td>
<td>8982/3</td>
</tr>
<tr>
<td>Carcinoma ex pleomorphic adenoma</td>
<td>8941/3</td>
</tr>
</tbody>
</table>

Acinic cell carcinoma

These are uncommon in minor glands (9, 280,340,410,734,864,2886) and form 2-6.5% of all intraoral salivary gland tumours (669,704,2711). In one series, the age range was from 11-77 years, with a mean of 45 years, and a male to female ratio of 1.5:1 (2711). The most common sites are the buccal mucosa, upper lip and palate where the tumours usually form non-descript swellings. The microscopic features are similar to those in major glands. However, in one series there appeared to be more areas consisting of solid sheets of epithelium with secretory material and fewer areas showing the minor glands (2711).

Mucoepidermoid carcinoma

This most common malignant salivary gland tumour involves minor glands, and accounts for 9.5-23% of all minor gland tumours (669,704,2711). About half of the cases arise in the palate and other common sites include the buccal mucosa, lips, floor of oral cavity and retromolar pad. They appear to be much more frequent in the lower lip than the upper lip (1871).

The tumour is often asymptomatic and detected during a routine dental examination. Many appear as bluish, domed swellings that resemble mucoceles or haemangiomas. Less commonly, the surface appears granular or papillary. Tumours of the base of tongue or oropharynx may cause dysphagia and sublingual tumours can lead to ankyloglossia and dysphonia. High-grade tumours are uncommon but can result in ulceration, loosening of teeth, parasthesia or anaesthesia. Mucoepidermoid carcinoma is the most common salivary gland tumour to develop in a central location within the bone of the mandible or, less frequently, the maxilla (280).

The microscopical features of minor gland mucoepidermoid carcinomas are the same as those seen in the major glands.

Adenoid cystic carcinoma

This lesion is relatively common in the minor glands. In the AFIP series 42.5% of all adenoid cystic carcinomas were in minor glands and 20.5% of the total was

Table 4.04 Percentage of malignant minor salivary gland tumours in different sites in published series.

<table>
<thead>
<tr>
<th></th>
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<tbody>
<tr>
<td>Upper Lip</td>
<td>22.5</td>
<td>14</td>
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</tr>
<tr>
<td>Lower Lip</td>
<td>60.2</td>
<td>86</td>
<td>-</td>
</tr>
<tr>
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<td>17.2</td>
<td>-</td>
<td>14.8</td>
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<tr>
<td>Palate</td>
<td>46.8</td>
<td>42</td>
<td>-</td>
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<tr>
<td>Tongue</td>
<td>85.7</td>
<td>-</td>
<td>47</td>
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<tr>
<td>Cheek</td>
<td>50.5</td>
<td>46</td>
<td>91.7</td>
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<td>Retro-molar</td>
<td>89.7</td>
<td>91</td>
<td>33.3</td>
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<tr>
<td>FOM*</td>
<td>88.2</td>
<td>80</td>
<td>-</td>
</tr>
<tr>
<td>Tonsil</td>
<td>41.4</td>
<td>-</td>
<td>50</td>
</tr>
<tr>
<td>Other</td>
<td>65.6</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

* Floor of the mouth
the tongue, tonsil and oropharynx, cheek, lips, retromolar pad and gingiva (899). They are much more frequent in the upper lip than the lower lip (669,1871,2711). Intraoral adenoid cystic carcinomas usually present as slow growing submucosal masses and ulceration may be seen, particularly in the palate. Pain, or evidence of nerve involvement, is usually only present in advanced tumours. Most tumours show the typical cylindromatous or cribriform variant microscopically, but some may have tubular areas and a few are predominantly solid (2711).

Polymorphous low-grade adenocarcinoma
This tumour is seen almost exclusively in minor glands and is considered in detail in Chapter 5.

Epithelial-myoepithelial carcinoma
This tumour is rare in minor glands and the literature consists mainly of single cases or short series (154,436,493,784, 981,992,1177). Tumours of the oral cavity and oropharynx formed only 10.3% of the AFIP series (669). The palate is the most common site. The clinical presentation is non-specific and the microscopical features are the same as those in major glands.

Basal cell adenocarcinoma
This tumour is rare in minor glands. There were none in the AFIP series (669) but there have been isolated case reports [563,785,1211,1540,2059,2703]. The most common sites are the palate, buccal mucosa and lip. They usually form asymptomatic, smooth or lobulated submucosal masses apart from one case that presented with dull pain and inflamed overlying mucosa (2703). Microscopically they are similar to basal cell adenocarcinomas of the major glands.

Cystadenocarcinoma
These tumours are uncommon and about 32% developed in the minor glands where they are frequently papillary (411,790). The most frequent sites are the palate, lips, buccal mucosa, tongue (1834) and retromolar regions. They are usually slow growing and painless but some palatal tumours have eroded the underlying bone and invaded the sinonasal complex. This tumour is considered in detail in Chapter 5.

Oncocytic carcinoma
This tumour is rare and there were only two cases involving the oral cavity in the AFIP series of 26 cases (669). One was in the palate and the other the buccal mucosa. Reported cases also include an additional case in the palate (274) and the AFIP case from the buccal mucosa (922). The microscopical features are considered in Chapter 5.

Salivary duct carcinoma
This tumour is rare in minor salivary glands. A recent review documented 20 cases (1147) and a further 6 cases have been reported (1559,2673). The most common location was the palate (65%). Other sites included the buccal mucosa and vestibule (19%), tongue (8%), retromolar pad (4%) and upper lip (4%). The age range was 23-80 years (mean 56 years). Some tumours formed painless swellings but many in the palate were painful and ulcerated or fungated. There were metastases to regional lymph nodes in 25% of cases and this was associated with a poor prognosis. The range of microscopical appearances was similar to that seen in the major glands.

Myoepithelial carcinoma
This is a rare salivary gland tumour and 26% of cases in a review of the literature (9 cases) involved the oral cavity or oropharynx (668) and only isolated cases have been published since this review (1827). The most common location is the palate. The clinical signs and symptoms are non-specific and the microscopical features are considered in Chapter 5.

Carcinoma ex pleomorphic adenoma
Lesions involving the oral and oropharyngeal minor glands formed 17.5% of the AFIP series (669). 63% of cases were in the palate and 10.5% were in the upper lip. There were no cases in the lower lip. Other sites included the tongue, buccal

---

**Fig. 4.32** A Mucoepidermoid carcinoma. Low power showing low-grade tumour with both cystic and solid areas and an inflamed, fibrous stroma. B Adenoid cystic carcinoma. This predominantly solid variant shows peri- and intraneural invasion. C Salivary duct carcinoma with large, somewhat oncocytic cells, cribriform areas, small papillae and comedo-type necrosis.
mucosa and tonsil/oropharynx. They usually form a painless mass of long duration and there may be a history of recent rapid growth, often with ulceration. The microscopical features are similar to those of major glands.

*Mucinous adenocarcinoma* is very rare while *clear cell carcinoma* is a controversial entity; both are discussed in Chapter 5.

**Salivary gland adenomas**

**ICD-O codes**

- Pleomorphic adenoma 8940/0
- Myoepithelioma 8982/0
- Basal cell adenoma 8147/0
- Canalicular adenoma 8149/0
- Duct papilloma 8503/0
- Cystadenoma 8440/0

*Pleomorphic adenoma*

These amount to 40-70% of minor gland tumours, the large majority of cases being located in the palate, lips and buccal mucosa [2711]. They usually present as painless, slow-growing, submucosal masses, but occasionally they are traumatised and bleed or ulcerate. They rarely exceed 3 sphere cm in diameter. Oral pleomorphic adenomas are similar microscopically to tumours elsewhere but frequently lack encapsulation, especially in the palate. They tend to be cellular, and hyaline or plasmacytoid cell types are common. Squamous metaplasia is also frequently seen and may be extensive. Some tumours have a strikingly lipomatous stroma and this should not be misinterpreted as tumour invading fat. Cases of intraoral pleomorphic adenoma with florid pseudoepitheliomatous hyperplasia of the overlying mucosa have been reported following incisional biopsy [2541].

*Myoepithelioma*

The minor glands are the common site for and myoepitheliomas account for about 42% of all of these tumours. Two thirds of the intraoral cases involve the palate [899]. They show the same range of morphological variation described in Chapter 5, but predominantly plasmacytoid tumours have a predilection for the palate of younger individuals [546].

*Basal cell adenoma*

About 20% of basal cell adenomas involve the oral cavity and the upper lip and buccal mucosa are the most common sites [669]. They are histologically similar to those in major glands.

*Cystadenoma*

These lesions are uncommon and form 7% of benign minor gland tumours [668]. Of these, 30% arose in the lips, 23% in the cheek, 20% in the palate and 26% in other oral and oropharyngeal sites. Clinically they resemble mucoceles and rarely exceed 1 cm in diameter. The pathology is discussed in Chapter 5.

*Canalicular adenoma and duct papillomas* arise almost exclusively in the minor salivary glands and are discussed in detail in Chapter 5.
Kaposi sarcoma

Definition
Kaposi sarcoma (KS) is a locally aggressive tumour that typically presents with cutaneous lesions in the form of multiple patches, plaques or nodules but may also involve mucosal sites, lymph nodes and visceral organs. The disease is uniformly associated with human herpes virus 8 (HHV-8) infection. KS rarely metastasizes and belongs to the group of intermediate type vascular tumours.

ICD-O code 9140/3

Epidemiology
Four different clinical and epidemiological forms of KS are recognized: 1. classic indolent form occurring predominantly in elderly men of Mediterranean/East European descent, 2. endemic African KS that occurs in middle-aged adults and children in Equatorial Africa who are not HIV infected, 3. iatrogenic KS appearing in solid organ transplant recipients treated with immunosuppressive therapy and also in patients treated by immunosuppressive agents, notably corticosteroids, for various diseases (2629), 4. acquired immunodeficiency syndrome-associated KS (AIDS KS), the most aggressive form of the disease, found in HIV-1 infected individuals, that is particularly frequent in homo- and bisexual men. The relative risk of acquiring KS in the latter patients is > 10,000 (909); it has been reduced with the advent of highly active antiretroviral therapy (HAART) (212), although this has not been proven yet for oral KS (1993,2120).

Etiology
The disease is the result of a complex interplay of HHV-8 with immunologic, genetic, and environmental factors (392). Oral exposure to infectious saliva seems to be a potential risk factor for the acquisition of HHV-8 (1995). HHV-8 is found in KS cells of all epidemiological-clinical forms of the disease (2242) and is detected in the peripheral blood before the development of KS. Nevertheless, it has been observed that a decline in incidence of KS did not appear to be caused by a decline in HHV-8 transmission (1959).

Localization
The most typical site of involvement by KS is the skin, particularly of the face and lower extremities. During the course of the disease or initially, mucosal membranes such as oral mucosa, lymph nodes and visceral organs may be affected, sometimes without skin involvement. Oral KS most frequently occurs on the palate, followed by the gingiva and the tongue.

Clinical features
Classic type of KS is characterized by the appearance of purplish, reddish blue or dark brown macules, plaques and nodules that may ulcerate. They are particularly frequent in distal extremities and may be accompanied by lymphoedema. The disease is usually indolent, lymph node and visceral involvement occurs infrequently. Classic KS may be associated with haematolymphoid malignancies.

In the endemic form of KS, the disease may be localized to skin and shows a protracted course. A variant of endemic disease, a lymphadenopathic form in African children is rapidly progressive and highly lethal.

Iatrogenic KS is relatively frequent. It develops in a few months to several years after the transplantation of solid organs or immunosuppressive treatment for a vari-

<table>
<thead>
<tr>
<th>Type</th>
<th>Risk groups</th>
<th>Skin lesions -predilection sites</th>
<th>Visceral involvement</th>
<th>Course</th>
</tr>
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<tbody>
<tr>
<td>Classic</td>
<td>Elderly men of Mediterranean/East European descent</td>
<td>Lower legs</td>
<td>Rare</td>
<td>Indolent</td>
</tr>
<tr>
<td>Endemic</td>
<td>Middles-aged men and children in Equatorial Africa</td>
<td>Extremities</td>
<td>Fairly common - adults</td>
<td>Indolent - adults</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Frequent - children (lymph nodes)</td>
<td>Aggressive - children</td>
</tr>
<tr>
<td>Iatrogenic</td>
<td>Immunosuppressed patients (post-transplant, other diseases)</td>
<td>Lower legs</td>
<td>Fairly common</td>
<td>Indolent or aggressive</td>
</tr>
<tr>
<td>AIDS-associated</td>
<td>Younger, mainly homo- and bisexual HIV-1 infected men</td>
<td>Face, genitalia, lower extremities</td>
<td>Frequent</td>
<td>Aggressive</td>
</tr>
</tbody>
</table>

From: WHO Classification of Tumours of Soft Tissue and Bone (175).
ety of conditions. The disease may resolve entirely upon withdrawal of immunosuppressive treatment although its course is somewhat unpredictable. AIDS-related KS is the most aggressive type of KS. Early oral KS is represented by solitary or multiple red or bluish flat lesions, while the later stage is characterized by a nodular, sometimes massive appearance with or without secondary ulceration.

**Histopathology**

Microscopic features of all four different epidemiological-clinical types of KS do not differ. Early lesions of the skin or the mucosa are uncharacteristic and present with subtle vascular proliferation [2216]. In the patch stage, vascular spaces are increased in number, of irregular shape, and may dissect collagen fibres in the superficial corium. They often run parallel to the epithelium. The vascular proliferation is often perivascular and periannexal. Endothelial cells lining the spaces are flattened or more oval, with little atypia. Pre-existing blood vessels may protrude into the lumen of new vessels. Admixed are sparse lymphocytes and plasma cells; frequently, extravasated erythrocytes and deposits of hemosiderin surround the vascular structures. Slits lined by attenuated endothelial cells between collagen bundles are also seen. In some cases, there is a proliferation of spindle or oval endothelial cells around pre-existing blood vessels in the dermis or submucosa. Slit-like spaces, lymphocyte and plasma cell infiltration and extravasated erythrocytes are also observed. In plaque stage, all characteristics of patch stage are exaggerated. There is more extensive angio-proliferation with vascular spaces showing jagged outlines. Inflammatory infiltrate is denser and extravascular red cells and siderophages are numerous. Hyaline globules (likely representing destroyed red blood cells) are frequently found. Nodular stage is characterized by well-defined nodules of intersecting fascicles of spindle cells with only mild atypia and numerous slit-like spaces containing red cells. Peripherally, there are ectatic blood vessels. Many spindle cells show mitoses. Hyaline globules are present inside and outside the spindle cells. Some patients, usually with endemic nodular type KS, develop lesions that closely resemble lymphangioma. The main differential diagnosis includes Kaposiform haemangioendothelioma [775].

**Immunoprofile**

The lining cells of clearly developed vascular structures are usually positive for vascular markers, while the spindle cells consistently show positive reaction for CD34 and commonly for CD31 but are factor VIII negative. All cases, irrespective of epidemiologic subgroup, are HHV-8 positive. The new marker FLI1, a nuclear transcription factor, appears to be expressed in almost 100% of different vascular tumours, including KS [780].

**Prognosis and predictive factors**

The evolution of disease depends on the epidemiological-clinical type of KS and on its clinical extent. It is also modified by treatment that includes surgery, radio- and chemotherapy. Patients with oral KS who did not receive triple antiretroviral therapy had a higher death rate than those having exclusively cutaneous manifestations of the disease [2192].
Lymphangioma

Definition
A benign, cavernous/cystic vascular lesion composed of dilated lymphatic channels.

ICD-O code 9170/0

Epidemiology
Lymphangiomas are common paediatric lesions, which most often present at birth or during the first years of life. Lymphangiomas appear mostly in the head and neck area but may be found in any other part of the body.

Etiology
Early or even congenital appearance in life and lesional architecture are in favour of a developmental malformation, with genetic abnormalities playing an additional role in cystic lymphangioma of the neck in association with Turner syndrome [416].

Clinical features
The lesion presents as a somewhat circumscribed painless swelling, which is soft and fluctuant on palpation. In oral involvement, the tongue is the site of predilection, the majority of lymphangiomas being located on the dorsal surface of the anterior part of the tongue. The size may vary from pinhead dimensions to massive lesions involving the entire tongue and surrounding structures. The typical lymphangioma of the tongue is characterized by irregular nodularity of the dorsum of the tongue with grey and pink, grapelike projections. Secondary haemorrhage in lymphangiomas is not a rare occurrence. CT scan reveals homogeneous non-enhancing areas [775]. A staging system of lymphatic malformations of the head and neck based on the anatomic location has shown to be of relevance in predicting prognosis and outcome of surgical intervention [561,991].

Macroscopy
Lymphangiomas form a multicystic or spongy mass, the cavities of which contain watery to milky fluid.

Histopathology
Lymphangiomas are characterized by thin-walled, dilated lymphatic vessels of different size, which are lined by a flattened endothelium. There is no encapsulation. The lumina may be either empty or contain proteinaceous fluid, lymphocytes and sometimes a few erythrocytes. Longstanding lesions show interstitial fibrosis.

Immunoprofile
The endothelium demonstrates variable expression of FVIII-rAg, CD31 and CD34 [781].

Electron microscopy
The endothelium of thin-walled vessels is not enveloped by a basement membrane and no pericytes are attached to it, thus directly contacting with the interstitium. With increasing calibre the vessels may acquire pericytes and smooth muscle, respectively.

Prognosis and predictive factors
Recurrences are due to incomplete removal. Current interest is centred on treating these lesions with sclerosing agents [2117], interferon [1953] or bleomycin [2903A]. Malignant transformation does not occur. There is an exceedingly rare case report of a squamous cell carcinoma arising in a lymphangioma of the tongue [203].

Fig. 4.37 Lymphangioma on the dorsum of the tongue.

Fig. 4.38 Lymphangioma. A Cavernous lymphangioma. B Flattened endothelial lining in lymphangioma.
Ectomesenchymal chondromyxoid tumour of the anterior tongue

Definition
A benign ectomesenchymal chondromyxoid tumour that arises in the anterior tongue.

Epidemiology
In 1995, nineteen cases of the previously undescribed entity were reported. Ever since, a few additional case reports have been published (1169). The reported age range varies from 9-78 years; there is no distinct sex predilection.

Clinical features
Most tumours presented as an otherwise asymptomatic, slow growing solitary nodule in the anterior dorsal tongue. The consistency may vary from firm to soft elastic.

Macroscopy
The cut surface has a gelatinous consistency with occasional foci of haemorrhage.

Histopathology
The tumour is usually well-circumscribed, but not encapsulated. Occasionally, muscle fibres and nerve branches may be entrapped within the tumour. It is composed of round, cup-shaped, fusiform, or polygonal cells with uniform small nuclei and moderate amounts of faintly basophilic cytoplasm; some tumours may show nuclear pleomorphism, hyperchromatism, and multinucleation, while mitotic figures are scarce (2410). In addition, the presence of myxoglobulosis-like changes has been reported (1169). Alcian blue stains at pH 0.4 and 2.5 are positive, while mucicarmine is usually faintly positive in the extracellular matrix. The tumour cells do not stain with the periodic acid-Shiff (PAS). In the histological differential diagnosis other myxoid and chondroid lesions should be excluded, such as focal oral mucinosis, the mucous retention phenomenon, soft-tissue myxoma, nerve sheath myxoma, myxomatous changes in fibrous lesions, chondrosarcoma, chondroid choriostoma, and variants of pleomorphic adenoma or myoepithelioma arising from minor salivary glands.

Immunoprofile
Reactivity with polyclonal and monoclonal anti glial fibrillary acidic protein (GFAP) is positive in almost all reported cases; reactivity with anti-cytokeratin monoclonal antibody has been positive in the majority of cases as well, while variable staining results were observed for S-100, CD57 and smooth muscle actin (2410).

Histogenesis
The tumour cells are possibly derived from undifferentiated ectomesenchymal progenitor cells that have migrated from the neural crest (2410).

Prognosis and predictive factors
Surgical excision is the treatment of choice. The recurrence rate is apparently low.

Fig. 4.39 Ectomesenchymal chondromyxoid tumour of the anterior tongue presenting as a small nodule.

Fig. 4.40 A Rather well demarcated ectomesenchymal chondromyxoid tumour. B Net-like pattern of round or ovoid cells in a chondromyxoid background.
**Focal oral mucinosis**

**Definition**
Focal oral mucinosis (FOM) is the oral counterpart of focal cutaneous mucinosis and cutaneous myxoid cyst. It is postulated that FOM develops as the result of a fibroblastic overproduction of hyaluronic acid due to an unknown cause (2615).

**Epidemiology**
Today, fewer than fifty cases have been reported. The lesion may occur at all ages, but it is rare in children (906). There is no distinct sex predilection.

**Clinical features**
The clinical presentation is usually that of an otherwise asymptomatic fibrous or cystic-like lesion. The most common site is the gingiva; less common sites include the palate, cheek mucosa and tongue. The consistency may vary from soft elastic to firm.

**Histopathology**
The histopathology is characterized by a well-circumscribed area of myxomatous tissue in which fusiform or stellate fibroblasts are present (299). Reticular fibres are sparse or absent. The mucinous material shows alcianophilia at pH 2.5. The histologic differential diagnosis includes soft-tissue myxoma, myxomatous change in fibrous lesions, nerve sheath myxoma, and mucous retention phenomenon. The lack of reticular fibres and the sharp delineation distinguishes FOM both from soft-tissue myxoma and from myxomatous changes in fibrous lesions (2615). Nerve sheath myxoma usually shows a lobular architecture and, conspicuously, contains numerous mast cells. The mucous retention phenomenon is surrounded by a wall of granulation tissue or an epithelium-lined wall, while the mucoid material contains histiocytic cells; such features are lacking in FOM.

**Prognosis and predictive factors**
The lesion is treated by conservative surgical excision and has no tendency to recur.
Congenital granular cell epulis

Definition
A benign tumour arising from the alveolar ridges of newborns and composed of nests of cells with granular cytoplasm set in a prominent vasculature [2744].

Synonym
Congenital epulis of the newborn

Epidemiology
In a review of the literature 216 cases have been collected since its first description in 1871. Females are affected ten times more often than males [2152].

Clinical features
Congenital granular cell epulis (CGCE) occurs twice as often in the maxilla as in the mandible, usually presenting as a solitary, somewhat pedunculated fibroma-like lesion attached to the alveolar ridge near the midline. A few cases of simultaneous occurrence of a CGCE and a granular cell tumour of the tongue have been reported [1564,2848]. The size of a CGCE may vary from a few millimetres up to several centimetres. Since the availability of ultrasound examination techniques, a number of cases have been diagnosed in the prenatal stage [1839,2000].

Histopathology
CGCE consists of large, slightly eosinophilic cells with granular cytoplasm set in a prominent vasculature. There is no cellular or nuclear pleomorphism, and mitotic activity is not usually observed. The presence of odontogenic epithelium scattered throughout the lesions has been reported. Immunohistochemically, the tumour cells are positive for vimentin and neuron specific enolase; there is no reactivity with cytokeratin, CEA, desmin, hormone receptors or S-100 [1968]. Pseudopitheliomatous changes in the overlying epithelium, although common in the granular cell tumour, do not occur in CGCE. An extremely rare case of a congenital leiomyomatous epulis has been reported [2542].

Histogenesis
The histogenesis is unknown. The lack of immunoreactivity with S-100 protein suggests that the tumour is derived from a cell line different from granular cell tumour. Furthermore, the hypothesis of a non-neoplastic lesion can be raised.

Prognosis and predictive factors
Spontaneous regression may occur, but surgical removal is usually indicated due to interference with feeding or respiration. The tumour has no tendency to recur after surgery.

Fig. 4.43 Congenital granular cell epulis of the maxilla.

Fig. 4.44 Congenital granular cell epulis. A Slightly eosinophilic cells with granular cytoplasm in a prominent vasculature. B Absence of nuclear and cellular pleomorphism.
Non-Hodgkin lymphoma

**Definition**
Non-Hodgkin lymphomas (NHL) of the oral cavity and oropharynx are defined as lymphoid cell neoplasms in which the bulk of the disease occurs in the palate, tongue, floor of mouth, gingiva, buccal mucosa, lips, palatine tonsils, lingual tonsils or oropharynx.

**ICD-O codes**
- Diffuse large B-cell lymphoma (DLBCL) 9680/3
- Mantle cell lymphoma 9673/3
- Follicular lymphoma 9690/3
- Extranodal marginal zone B-cell lymphoma of MALT type 9699/3
- Burkitt lymphoma 9687/3
- T-cell lymphoma (including anaplastic large cell lymphoma) 9714/3

**Epidemiology**
Although NHL is the second most common cancer of the oral cavity, it only accounts for 3.5% of all oral malignancies [683]. NHL of the oral cavity and oropharynx account for 13% of all primary extranodal NHL, with approximately 70% of these occurring in the tonsils [809]. They affect patients over a wide age range (including children), but most patients are in the 6th and 7th decades. Burkitt lymphomas occur predominantly in children and young adults. Patients with an underlying immunodeficiency state (e.g. HIV Infection) are also usually younger.

**Etiology**
There is no known etiology in most patients. A minority of patients have an underlying immunodeficiency state (e.g. HIV infection, post-transplantation), which predisposes to the development of NHL. There is a strong association with Epstein-Barr virus (EBV) for lymphomas occurring in the setting of immunodeficiency as well as in extranodal NK/T cell lymphoma of nasal-type [371,962,1476,2850]. Extranodal marginal zone B-cell lymphoma of MALT type may be associated with Sjögren syndrome [2476].

**Localization**
The palatine tonsil is the most frequently involved site, followed by palate, gingiva and tongue [683,809,1476,2532].

**Clinical features**
Patients with NHL of the lip, buccal mucosa, gingiva, floor of mouth, tongue or palate usually present with ulcer, swelling, discoloration, pain, paraesthesia, anaesthesia, or loose teeth. Those with NHL of the Waldeyer ring (tonsils) or oropharynx usually present with a sensation of fullness of the throat, sore throat, dysphagia, or snoring. The high-grade tumours often show rapid growth. Systemic symptoms such as fever and night sweat are uncommon [201]. Clinical examination reveals solitary or multiple lesions, in the form of an exophytic mass, ulcer or localized swelling. Some cases may mimic inflammatory

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Table 4.06 Frequency of the various types of primary lymphoma of the tonsil (1704)

<table>
<thead>
<tr>
<th>Lymphoma Type</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diffuse large B-cell lymphoma (DLBCL)</td>
<td>64%</td>
</tr>
<tr>
<td>Mantle cell lymphoma</td>
<td>10%</td>
</tr>
<tr>
<td>Follicular lymphoma</td>
<td>8%</td>
</tr>
<tr>
<td>Extranodal marginal zone B-cell lymphoma of MALT type*</td>
<td>6.5%</td>
</tr>
<tr>
<td>T-cell lymphoma (including anaplastic large cell lymphoma)</td>
<td>6.5%</td>
</tr>
<tr>
<td>Burkitt lymphoma</td>
<td>5%</td>
</tr>
</tbody>
</table>

* Reported as monocytoid B-cell lymphoma and immunocytoma in the original series.

---

Fig. 4.45 A Primary large B cell lymphoma of the lip. The mucosa shows a diffuse infiltrate of lymphoma cells beneath an intact stratified squamous epithelium. B Primary large B cell lymphoma of the tongue. In this example, the mucosa is partly ulcerated.
conditions, such as periodontitis. Tonsillar lymphoma usually manifests as asymmetric tonsil enlargement, although the disease can be bilateral in up to 9% of cases [2250]. The regional lymph nodes can be enlarged as a result of lymphoma involvement or reactive changes secondary to ulceration.

**Tumour spread and staging**

Three-quarters of patients have localized disease, with or without accompanying cervical lymph node involvement at presentation (Stage IE/IIE). Patients with lymphoma of the tonsil are prone to metachronous or synchronous gastrointestinal tract involvement, suggesting a homing mechanism among different mucosal sites [1998,2139,2250].

**Histopathology**

Most NHL of the oral cavity and oropharynx are of B-cell lineage, with DLBCL being the commonest (>50%) [370,1476,1704,2142,2530]. The surface stratified squamous epithelium is either intact or ulcerated. The stroma is densely infiltrated by lymphoma cells, which vary in appearance depending on the histologic type. In the tonsil, not uncommonly there are some residual lymphoid follicles due to incomplete involvement of the tissue.

**Diffuse large B-cell lymphoma (DLBCL)**

DLBCL is characterized by large to medium-sized cells which may resemble centroblasts. Nuclear multilobation is prominent in some cases. There can be areas of coagulative necrosis. In DLBCL of the tonsils, a focal follicular pattern may be present [2138], and it has been argued that the follicles result from colonization of pre-existing follicles rather than de novo neoplastic follicle formation. In some cases, there may be an associated component of extranodal marginal zone B-cell lymphoma of MALT type or follicular lymphoma, indicating that the DLBCL represents high-grade transformation of the latter [1998].

**Mantle cell lymphoma**

Lymphoma cells are usually monotonous, and frequently have small irregular nuclei, dense chromatin and scanty cytoplasm. They may show a mantle zone distribution around residual follicles. Rare cases can have a blastic appearance and are associated with a higher proliferation rate [19].

**Follicular lymphoma**

Follicular lymphoma is characterized by follicles that frequently lack polarity and mantle zone. The neoplastic follicles consist of a mixture of centrocytes and centroblasts, often without accompanying tingible-body macrophages.

**Extranodal marginal zone B-cell lymphomas of MALT type**

These lymphomas most often involve the tonsil, and less commonly the palate, gingiva, buccal mucosa, tongue and lip [295,962,1476,1507,1629,1998,2225,2531]. The surface epithelium is often intact. In a background of reactive lymphoid follicles, there is an interfollicular and perifollicular infiltrate of small to medium-sized cells with roundish or indented nuclei. Some cells have a moderate amount of clear cytoplasm, resembling monocytoid B-cells. There can be clusters of admixed plasma cells. Follicular colonization can be seen in some cases. A distinctive feature is invasion of the epithelial component (e.g. sur-
face or crypt epithelium, minor salivary glands), forming lymphoepithelial lesions.

**Burkitt lymphoma**

There is typically a starry sky pattern created by interspersed histiocytes. The lymphoma cells appear monotonous and medium-sized, with coarse chromatin, multiple small nucleoli and a small amount of basophilic cytoplasm. The cellular outline usually appears squared off. Frequent mitotic figures and apoptotic bodies are constant features.

**Immunoprofile.** B-cell lymphomas express pan-B markers such as CD19, CD20, CD22 and CD79a. Some DLBCLs can express CD10 and/or BCL6. Within the group of low-grade B-cell lymphomas, follicular lymphoma is characterized by CD10 and BCL6 expression, mantle cell lymphoma CD5 and cyclin D1 expression, and extranodal marginal zone B-cell lymphoma none of these markers. Bcl-2 Immunoreactivity is helpful for distinction of follicular lymphoma from reactive follicular hyperplasia.

**Extranodal NK/T cell lymphoma of nasal-type**

Extranodal NK/T cell lymphoma of nasal-type can present primarily as an intraoral tumour in the palate, tonsil, oropharynx or lip [371, 2639]. Please refer to the section of ‘non-Hodgkin lymphoma’ in ‘Tumours of the nasal cavity and paranasal sinuses’ for details.

**T-cell lymphomas**

Peripheral T-cell lymphomas, including anaplastic large cell lymphomas, can occasionally involve the oral cavity [1476, 2200, 2530]. Some anaplastic large cell lymphomas (CD30+ T-cell lymphoproliferative disorder) of the oral cavity can regress spontaneously [760]. HTLV-1-associated adult T-cell lymphoma/leukaemia may also present as NHL of the Waldeyer ring [2616].

**Immunodeficiency-associated lymphomas**

The lymphomas that develop in the oral cavity of patients with HIV infection are most commonly DLBCL with frequent EBV association (75%) [962, 2141], although EBV-associated T-cell lymphomas have also been reported in this setting [1476, 2589]. A distinctive form of DLBCL, plasmablastic lymphoma, has recently been shown to exhibit a predilection for the oral cavity of HIV-positive subjects. It differs from the usual
Tumours of the oral cavity and oropharynx

**Differential diagnoses**

In *infectious mononucleosis*, the tonsils may appear histologically worrisome, with necrosis, partial effacement of architecture, and striking immunoblastic proliferation, mimicking DLBCL (431). In contrast to the latter, there is usually a spectrum of lymphoid cells in different stages of differentiation and activation (immunoblasts, plasmablasts and plasma cells). On immunostaining, the large cells usually consist of a mixture of B- and T-cells, and there is no immunoglobulin light chain restriction. As a rule of thumb, infectious mononucleosis has to be seriously excluded before making a diagnosis of DLBCL in young patients.

Some cases of DLBCL (especially those in the tonsil) can exhibit deceptively cohesive growth and a sharp interface with the uninvolved mucosa, closely mimicking poorly differentiated carcinoma or malignant melanoma. Marked irregular nuclear foldings and amphophilic cytoplasm, if present, should point more towards a diagnosis of lymphoma. Appropriate immunostains can readily solve this diagnostic problem.

Anaplastic plasmacytoma can be difficult to distinguish from DLBCL, including the plasmablastic variant. An important clue to the diagnosis is the presence of coarsely clumped ‘clock-face’ chromatin in the few differentiated cells that are present. There are often intermingled atypical plasma cells. There is usually no association with EBV. A prior history of multiple myeloma, if present, would be a strong point to substantiate a diagnosis of plasmacytoma.

Extramedullary myeloid sarcoma (granulocytic sarcoma) is commonly misdiagnosed as large cell lymphoma. The clues to diagnosis are the fine chromatin, presence of cytoplasmic eosinophilic granules in some cells, and interspersed eosinophilic myelocytes. The diagnosis can be confirmed by immunoreactivity for myeloid or monocytic markers (e.g. myeloperoxidase, CD13, CD33, CD117, neutrophil elastase, lysozyme, CD68).

The differential diagnosis between extranodal marginal zone B-cell lymphoma of MALT type in the tonsil and *reactive lymphoid hyperplasia* can be extremely difficult, because of the presence of reactive lymphoid follicles, minimal atypia of the lymphoid cells in the former and presence of numerous plasma cells. Furthermore, lymphoepithelial lesions in the tonsil are difficult to assess since the tonsillar epithelium is normally extensively infiltrated by small lymphoid cells. The following features would favour a diagnosis of lymphoma: lymphoid cells infiltrating beyond the fibrous band at the base of the tonsil, presence of sheets of CD20+ B-cells between the lymphoid follicles, immunoglobulin light chain restriction, and molecular evidence of clonal immunoglobulin gene rearrangement.

Some extranodal NK/T cell lymphomas of *nasal-type* comprise predominantly small lymphoid cells with minimal atypia, rendering it difficult to distinguish from a reactive lymphoid infiltrate. Histologic clues to the diagnosis are the extensive necrosis and angiocentric growth. Demonstration of sheets of CD56+ or EBER+ cells would strongly support the diagnosis.

There is some morphologic overlap of anaplastic large cell lymphoma with *eosinophilic ulcer* (traumatic eosinophilic granuloma; atypical histiocytic granuloma) (645,674,701), which is characterized by a rich inflammatory infiltrate (especially eosinophils) and occasional large cells (760). Anaplastic large cell lymphoma can be distinguished from it by the presence of at least large aggregates of large atypical cells in areas and strong CD30 expression.

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**Fig. 4.50 Plasmacytoma of the tongue.** The plasmacytoma is accompanied by blood lakes.
Haematolymphoid tumours

Prognosis and predictive factors

Patients with NHL of the oral cavity and oropharynx are treated by radiotherapy, chemotherapy or a combination of the two. Some studies have shown that adjuvant chemotherapy is associated with a better clinical outcome compared to radiotherapy alone [832, 1009]. The five-year overall survival rate for localized disease ranges from 50% to more than 80% [146, 832, 1009, 1614, 2505]. High clinical stage, high histologic grade (large cell lymphoma), and T-cell or NK/T cell phenotype are poor prognostic indicators [146, 1009, 2250, 2340, 2809].

Extramedullary plasmacytoma

ICD-O code 9734/3

Extramedullary plasmacytoma can occur in the oral cavity and oropharynx, see Chapter 1 for details.

Langerhans cell histiocytosis

ICD-O code 9751/1

Oral involvement occurs in 10% of patients with Langerhans cell histiocytosis (LCH). 78% of these patients have eosinophilic granulomas clinically, while the rest have multifocal multisystem disease [1021]. Common oral symptoms include swelling, pain, gingivitis, loose teeth and ulceration. The majority of patients with intraoral lesions have intraosseous lesions in the jaw bone, more commonly in the mandible. The intraoral soft tissues may be secondarily affected, especially the gingiva, but the palate, floor of mouth, buccal mucosa and tonsil can also be involved [1021, 2043]. In a minority of patients with intraoral soft tissue involvement, there is no associated bony lesion [241, 460, 1731]. See chapter 7 for details.

Hodgkin lymphoma

Hodgkin lymphoma (HL) is predominantly a nodal-based disease, and primary extranodal presentation is very rare. When it presents in extranodal tissues, the Waldeyer ring, particularly the palatine tonsil, is a common site [1274, 1756]. Most patients present with localized disease (stage I/II), with symptoms of chronic tonsillitis or tonsillar enlargement, with or without enlarged cervical lymph nodes. Other reported sites include the oropharynx [44, 1756], alveolar crest of mandible [1659], and maxillary gingiva [2554]. Most cases represent classical HL, as detailed in the WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues [1197], frequently of mixed cellularity subtype and showing strong association with Epstein-Barr virus (EBV) [1274], although nodular lymphocyte predominant HL may also rarely present in the Waldeyer ring (palatine and lingual tonsils) [391, 1274].

Extramedullary myeloid sarcoma

ICD-O code 9930/3

Gingival infiltrates occur in 3.5% of patients with acute myeloid leukaemia, predominantly in the monocytic or myelomonocytic subtypes [622]. Clinically, there is diffuse enlargement of the interdental papillae, marginal gingiva and attached gingiva. The swollen gingiva has a spongy to firm consistency, bright red to purple in colour. There is no correlation between gingival leukaemic infiltrate and oral hygiene or peripheral white blood cell count [622]. Rare cases of extramedullary myeloid sarcoma may present as an isolated tumour-forming intraoral mass. The most frequently involved sites are the palate and gingiva [52, 761, 2189, 2614, 2618]. While the tumour most often develops while the patient has active disease, it may precede the development of acute myeloid leukaemia, or arise as a result of a chronic myeloproliferative disease or myelodys-
plastic syndrome. Histologically, there is a dense infiltrate of immature myeloid cells in the subepithelial soft tissue of the gingiva. Please refer to the section of ‘Other uncommon haematolymphoid tumours’ in ‘Tumours of the nasal cavity and paranasal sinuses’ for further details on extramedullary myeloid sarcoma.

**Follicular dendritic cell sarcoma / tumour**

**Definition**
Follicular dendritic cell (FDC) sarcoma/tumour is a rare neoplasm showing morphologic and phenotypic features of FDC.

**ICD-O code**
9758/3

**Epidemiology, localization and clinical features**
It is an uncommon tumour of adulthood, and can affect patients over a wide age range (368,2010,2043). It can arise in nodal and extranodal tissues, and the oral cavity is among the more commonly involved extranodal sites (67,368,375, 2010,2043,2249). The patients usually present with a painless mass involving the tonsil, palate or oropharynx.

**Etiology**
Occasional FDC sarcomas/tumours appear to evolve from an underlying ha-line-vascular Castleman disease; the two lesions can present simultaneously or the latter can precede the appearance of the former by several years (359,368,374). Overgrowth of FDC in the interfollicular zone or ‘dysplasia’ of FDC may represent the precursor lesion.

**Histopathology**
Histologically, the tumour usually grows beneath an intact stratified squamous epithelium. It usually exhibits pushing borders and comprises fascicles, whorls, nodules, storiform arrays or diffuse sheets of spindly to ovoid tumour cells sprinkled with small lymphocytes. The tumour cells usually show ill-defined cell borders, distinct nucleoli, and sometimes nuclear pseudoinclusions. There is a tendency for some nuclei to be haphazardly clustered, and scattered multinucleated tumour cells are common. While nuclear pleomorphism is usually mild, some cases can show significant nuclear atypia and pleomorphism. The cytoplasm is eosinophilic, and often exhibits a fibrillary quality as a result of the presence of interdigitating cell processes. Very rarely, the tumour cells have distinct cell borders, and are polygonal or oval in shape. The mitotic count ranges from low to high, and some cases may show coagulative necrosis. Occasional cases may show irregular interspersed cystic spaces. Besides being intermingled among the tumour cells, the lymphocytes can show cuffing around the blood vessels. The diagnosis has to be confirmed by demonstration of FDC markers (e.g. CD21, CD23 and CD35), although the staining can be patchy. Typically a mesh-
work pattern is highlighted. Cytokeratin is negative. A proportion of cases express epithelial membrane antigen or muscle-specific actin. Occasional cases can weakly express the pan-B marker CD20 (2043). Ultrastructurally, the tumour cells possess interdigitating long slender cytoplasmic processes and intercellular desmosome junctions. Differential diagnoses include soft tissue sarcoma, poorly differentiated carcinoma, meningioma, and malignant melanoma.

**Prognosis and predictive factors**
Most cases of FDC sarcomas/tumours have been treated by surgery, with or without adjuvant chemotherapy and radiotherapy, with variable success. FDC sarcomas/tumours are low to intermediate grade malignant tumours, with an overall local recurrence rate of at least 40% and a metastatic rate of at least 28% (368,2010,2043). Since some patients can develop late metastasis (such as after more than 20 years) (438), long-term follow up is essential. Poor prognostic factors include significant cytologic atypia, extensive coagulative necrosis, high proliferative index and large tumour size (368,2010).
Mucosal malignant melanoma

Definition
Malignant melanoma is a malignant neoplasm of melanocytes or of melanocyte precursors. It is characterized by proliferation of atypical melanocytes at the epithelial-connective tissue interface associated with upward migration into the epithelium and by invasion of the underlying connective tissues. Although usually seen in the skin, melanomas may also arise from melanocytes in mucosae.

ICD-O code 8720/3

Epidemiology
Mucosal melanomas of the head and neck comprise just over 1% of all melanomas and of these about 50% arise in the oral cavity. Oral mucosal melanomas are therefore rare, representing about 0.5% of all oral biopsies [122]. The annual incidence in the USA is about 0.02 per 100,000 [170], but the lesion may be more common in other parts of the world including Japan where the oral cavity has been reported as the most common site for melanomas [2528].

They arise in adults with an average age of about 55, but with a uniform age distribution from 20–80 years [122,617,1085,2080,2127]. Very rare cases have been reported in children. In most large series there is a male predominance in a ratio of about 3:1 [122,617,2080] and some reports show males and females are almost equal [144,1843].

Etiology
No etiological factors are known to be associated with oral melanoma.

Localization
Eighty percent of oral melanomas arise on the palate, maxillary alveolus or gingivae. Other sites include the mandibular gingivae, buccal mucosa, floor of mouth and tongue.

Clinical features
Oral melanomas are usually asymmetric with irregular outlines. They may be black, grey or purple to red, and rarely amelanotic. Typical lesions are composed of multiple or widespread areas of macular pigmentation with areas of nodular growth. Purely macular lesions may be seen but over 50% of lesions present as nodules or as a pigmented epulis. Ulceration is seen in about one third of cases and invasion of bone is common. Many reports document long-standing ‘melanosis’ before the onset of nodular lesions, with a history of up to 10 years. Oral lesions are usually advanced at presentation with up to 75% of patients having metastases to cervical lymph nodes, and 50% with distant metastases, usually to lung or liver [144,617,1085].

Macroscopy
Tumours are usually 1.5-4 cm in diameter with a black, macular or nodular surface. The cut surface is often homogeneously black or darkly pigmented.

Histopathology
Oral melanoma may have in-situ (radial) and invasive growth phases, but the histological classification is not analogous to cutaneous lesions. Mucosal lesions are similar to acral lentiginous melanoma of the skin [2652], with junctional activity and upward migration but Pagetoid invasion is unusual. Atypical melanocytic lesions may progress to malignant melanoma but there is little evidence for progression of oral benign melanocytic naevi to invasive malignancy [1085,2652]. Oral mucosal melanoma is, therefore, classified as in-situ oral mucosal melanoma, invasive oral mucosal melanoma, and mixed in-situ and invasive lesions. Borderline lesions may be termed atypical melanocytic proliferations [122,1085,2652]. Most lesions at presentation are invasive or have mixed invasive and in-situ com-

Fig. 4.56 Malignant melanoma. multiple or widespread areas of dark macular pigmentation affecting the palate. Irregular nodular areas are also seen.

Fig. 4.57 A in-situ growth phase showing atypical and enlarged melanocytes at the epithelial-connective tissue interface. Melanocytes may show upward migration into the epithelium. B Invasive lesions showing considerable junctional activity, with atypical melanocytes invading into the underlying connective tissues.
ponents. Less than 20% are solely in-situ lesions. Typically, an oral melanoma is composed of sheets or islands of epithelioid melanocytes, which may be arranged in an organoid, or alveolar pattern. The cells have pale cytoplasm and large open nuclei with prominent nucleoli and occasionally they may be plasmacytoid. Sheets and fascicles of spindle cells may also be seen, but are usually a minor part of the lesion. Occasional lesions may be predominantly or wholly spindle. Over 90% of lesions contain melanin pigment that can easily be demonstrated with stains such as Masson-Fontana or Schmorl's. When present, the in-situ component shows atypical naevoid cells arranged singly or in nests at the epithelial-connective tissue interface. Upward migration of the cells is common, but Pagetoid islands, similar to those of superficial spreading cutaneous melanomas, are not frequent. Invasion may be difficult to determine but the presence of obviously malignant cells in the lamina propria indicates invasion and islands of cells larger than those seen within the epithelium suggest an invasive growth phase. Mitoses are surprisingly sparse but are seen more frequently in invasive lesions. The overlying epithelium is usually atrophic and just over half of lesions are ulcerated.

**Immunoprofile**

Over 95% of lesions are S100 positive and negative for cytokeratins [144]. Although sensitive, S100 is not specific. More specific markers include HMB45, Melan-A or anti-tyrosinase, which stain about 75% of lesions [2079].

**Genetics**

Cutaneous melanomas may be associated with familial melanoma syndromes, and melanoma-prone kindreds show frequent loss of heterozygosity or mutations at several sites. Two tumour suppressor genes, CDKN2A (at 9p21, which codes for P16INK4A) and PTEN (at 10q23), and the oncogene CDK4 have been identified as important melanoma susceptibility genes [269,1252,2713]. However, associations with these genes have not yet been shown for oral melanomas, and expression of various tumour suppressor genes or oncogenes is variable and heterogeneous [1085,2555].

**Prognosis and predictive factors**

The prognosis for oral melanoma is poor with an overall median survival of about 2 years and 5-year survival of less than 20% [122,170,1085]. Stage is a predictor of survival but even localized tumours (stage I) show a 5-year survival of less than 50%. Depth of invasion (Breslow thickness and Clark's levels) is of limited value in oral lesions. This is due to lack of adequate studies and the fact that most oral melanomas are deeper than 4 mm at presentation [1085,1843,2080]. Nevertheless, lesions thicker than 5 mm may have a significantly worse prognosis. Other factors associated with poor prognosis include, vascular invasion, necrosis, a polymorphous tumour cell population, and increasing age [170,1085,1843,2080].
Metastases to bone

Definition
Distant spread of malignant neoplasm to the head and neck from other parts of the body. This is almost exclusively via a haematogenous route.

Epidemiology
The most common malignant neoplasms within the jaws, apart from direct spread from mucosal carcinomas, are metastases and the most frequent primary sites are carcinomas of, in order of decreasing incidence, breast, kidney, lung, prostate and thyroid or colon (1098). Maxillary and sinus metastases most frequently arise from renal carcinoma (202). Metastasis accounts for approximately 4% of all upper aerodigestive tract carcinoma (246). The great majority of patients are elderly, with mean age at diagnosis of 55 years and the sex incidence varies from equal (2877) to a female preponderance accounted for by the prevalence of breast carcinoma (202). Metastasis accounts for approximately 4% of all upper aerodigestive tract carcinoma (246). The great majority of patients are elderly, with mean age at diagnosis of 55 years and the sex incidence varies from equal (2877) to a female preponderance accounted for by the prevalence of breast carcinoma (202).

Clinical features
Common signs and symptoms include loosening of teeth, swelling, failure to heal of a dental extraction socket (1101), pathological fracture or nerve signs, particularly paraesthesia and anaesthesia in the mental region. Pain may be the only evidence of metastasis. After cortical perforation, a soft tissue mass may be present. Some metastases are asymptomatic chance radiographic findings.

Tumour spread and staging
Jaw metastases are the presenting sign of malignancy in 20-30% of cases (1098, 2877) but in most the primary lesion is known. Sometimes metastasis develops many years after treatment for the primary lesion, particularly with renal carcinoma. Metastatic spread to the jaws indicates UICC/AJCC Stage IV disease.

Histopathology
Histopathological appearances vary. Metastases are usually poorly-differentiated. If immunocytochemistry is required to aid clinical identification of sites of an occult primary lesion, prostate specific antigen and thyroglobulin are the most useful stains.

Prognosis and predictive factors
Metastasis to the jaws usually indicates widely disseminated disease and a poor prognosis with a 4-year survival of 10% (458). Two thirds of patients die in less than 1 year (1101). Depending on lesion type and dissemination, radiotherapy or hormone therapy may be provided. Surgery may occasionally be of value in palliative care.

Metastases to oral soft tissues
Metastasis to soft tissues is much more rare. It affects a similar age group, 40-70 years old, and the commonest sites for primary lesions in males are lung (one third of cases) followed by kidney and skin. The commonest primary site in females is breast (1100). The commonest site for metastasis is gingiva (55%) because of its fine capillary bed, followed by tongue (30%), though any site may be affected. The predilection for gingiva is mostly lost after teeth are extracted (1100). Lesions present as soft tissue masses, often ulcerated, resembling traumatic or reactive hyperplastic lesions.
CHAPTER 5

Tumours of the Salivary Glands

Salivary gland tumours can show a striking range of morphological diversity between different tumour types and sometimes within an individual tumour mass. In addition, hybrid tumours, dedifferentiation and the propensity for some benign tumours to progress to malignancy can confound histopathological interpretation. These features, together with the relative rarity of a number of tumours, can sometimes make diagnosis difficult, despite the abundance of named tumour entities. The increasing use of pre-operative fine needle aspiration biopsies also needs to be taken into account, as artifactual changes may be superimposed on the tumours. Unfortunately, the morphological variability of these tumours is mirrored by the immunocytochemical profiles, so that special stains are rarely useful in routine diagnosis of salivary gland epithelial neoplasms.
WHO histological classification of tumours of the salivary glands

<table>
<thead>
<tr>
<th>Malignant epithelial tumours</th>
<th>Basal cell adenoma</th>
<th>8147/0</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acinic cell carcinoma</td>
<td>Warthin tumour</td>
<td>8561/0</td>
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<tr>
<td>Mucoepidermoid carcinoma</td>
<td>Oncocytoma</td>
<td>8290/0</td>
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<tr>
<td>Adenoid cystic carcinoma</td>
<td>Canalicular adenoma</td>
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<tr>
<td>Polymorphous low-grade adenocarcinoma</td>
<td>Sebaceous adenoma</td>
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<td>Epithelial-myoepithelial carcinoma</td>
<td>Lymphadenoma</td>
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<td>Clear cell carcinoma, not otherwise specified</td>
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<td>8410/0</td>
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<td>Basal cell adenocarcinoma</td>
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<td>Ductal papillomas</td>
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<td>Adenocarcinoma, not otherwise specified</td>
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<td>Diffuse large B-cell lymphoma</td>
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<td>Extranodal marginal zone B-cell lymphoma</td>
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<tr>
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<td>Myoepithelioma</td>
<td>8892/0</td>
</tr>
</tbody>
</table>

1 Morphology code of the International Classification of Diseases for Oncology (ICD-O) (821) and the Systematized Nomenclature of Medicine (http://snomed.org). Behaviour is coded /0 for benign tumours, /3 for malignant tumours, and /1 for borderline or uncertain behaviour.
## TNM classification of carcinomas of the salivary glands

<table>
<thead>
<tr>
<th><strong>TNM classification</strong></th>
<th><strong>Note:</strong></th>
<th><strong>Stage Grouping</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>T – Primary tumour</strong></td>
<td><em>Extraparenchymal extension is clinical or macroscopic evidence of invasion of soft tissues or nerve, except those listed under T4a and 4b. Microscopic evidence alone does not constitute extraparenchymal extension for classification purposes.</em></td>
<td><strong>Stage I</strong> T1 N0 M0</td>
</tr>
<tr>
<td>TX</td>
<td>Primary tumour cannot be assessed</td>
<td><strong>Stage II</strong> T2 N0 M0</td>
</tr>
<tr>
<td>T0</td>
<td>No evidence of primary tumour</td>
<td><strong>Stage III</strong> T3 N0 M0</td>
</tr>
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<td>T1</td>
<td>Tumour 2 cm or less in greatest dimension without extraparenchymal extension*</td>
<td><strong>Stage IV A</strong> T1, T2, T3 N2 M0</td>
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<tr>
<td>T2</td>
<td>Tumour more than 2 cm but not more than 4 cm in greatest dimension without extraparenchymal extension*</td>
<td><strong>Stage IV B</strong> T4b Any N M0</td>
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<tr>
<td>T3</td>
<td>Tumour more than 4 cm and/or tumour with extraparenchymal extension*</td>
<td><strong>Stage IV C</strong> Any T Any N M1</td>
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<tr>
<td>T4a</td>
<td>Tumour invades skin, mandible, ear canal, or facial nerve</td>
<td><strong>Stage IV D</strong> Any T Any N M1</td>
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<tr>
<td>T4b</td>
<td>Tumour invades base of skull, pterygoid plates, or encases carotid artery</td>
<td><strong>Stage IV E</strong> Any T Any N M1</td>
</tr>
</tbody>
</table>

| | **Note:** | |
| | Midline nodes are considered ipsilateral nodes. | |
| | **N – Regional lymph nodes** | |
| | | **Stage I** T1 N0 M0 |
| | NX | Regional lymph nodes cannot be assessed | **Stage II** T2 N0 M0 |
| | N0 | No regional lymph node metastasis | **Stage III** T3 N0 M0 |
| | N1 | Metastasis in a single ipsilateral lymph node, 3 cm or less in greatest dimension | **Stage IV A** T1, T2, T3 N2 M0 |
| | N2 | Metastasis as specified in N2a, 2b, 2c below | **Stage IV B** T4b Any N M0 |
| | N2a | Metastasis in a single ipsilateral lymph node, more than 3 cm but not more than 6 cm in greatest dimension | **Stage IV C** Any T Any N M1 |
| | N2b | Metastasis in multiple ipsilateral lymph nodes, none more than 6 cm in greatest dimension | **Stage IV D** Any T Any N M1 |
| | | **Stage IV E** Any T Any N M1 |

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1 (947,2418).

2 A help desk for specific questions about the TNM classification is available at http://www.uicc.org/index.php?id=508.
Tumours of the salivary glands:
Introduction

Anatomy
Salivary glands are exocrine organs responsible for the production and secretion of saliva. They comprise the three paired major glands, the parotid, submandibular and sublingual, and the minor glands. The latter are numerous and are widely distributed throughout the mouth and oropharynx and similar glands are present in the upper respiratory and sinonasal tracts, and the paranasal sinuses.

Secretory acinus
The functional unit of salivary glands is the secretory acinus and related ducts, and myoepithelial cells. Acini may be serous, mucous or mixed. Serous acini form wedge-shaped secretory cells with basal nuclei. They surround a lumen that becomes the origin of the intercalated duct. The cytoplasm of serous cells contains densely basophilic, refractile zymogen granules that are periodic acid Schiff positive and diastase resistant. Their principle secretion is amylase. Mucous acinar cells also have basally placed nuclei and their cytoplasm is clear and contains vacuoles of sialomucin. The secretions of these cells pass through the intercalated ducts. These are often inconspicuous in routine histological sections. They are lined by what appears to be a single layer of cuboidal cells with relatively large, central nuclei. They are continuous with the much larger striated ducts. The intercalated ducts are lined by a single layer of cuboidal cells with relatively large, central nuclei and are linked to the much larger striated duct. The latter are lined by tall, columnar, eosinophilic cells that are rich in mitochondria. They have parallel infoldings of the basal cytoplasm and are responsible for modifying the salivary secretions. The striated ducts join the interlobular excretory ducts, which are lined by pseudostratified columnar epithelium that often contains few mucous cells.

Myoepithelial cells
Myoepithelial, or basket cells, are contractile and are located between the basement membrane and the basal plasma membrane of the acinar cells. They are variable in morphology and are inconspicuous in H&E sections. They contain smooth muscle actin, myosin and intermediate filaments including keratin 14. Immunohistochemical stains for the proteins highlights their stellate shape. They have long dendritic processes that embrace the secretory acini. Myoepithelial cells also surround the intercalated ducts but their presence in striated ducts is not firmly established. Ultrastructurally, the cytoplasm of myoepithelial cells contains actomyosin microfilaments running parallel with the outer surface of the cell, glycogen granules and lipofuscin, and pinocytotic vesicles may also be a conspicuous feature.

Parotid gland
The parotid gland is almost purely serous and the parenchyma is divided into lobules by fibrous septa. There is abundant intralobular and extralobular adipose tissue which increases in relative volume with age. The parotid gland contains randomly distributed lymphoid aggregates and lymph nodes that range from one to more than 20 in number. Not infrequently the lymph nodes contain salivary gland ducts or occasionally acini (Neisse Nicholson rests). Sebaceous glands, either individually or in small groups, are commonly seen if the tissue is widely sampled.

Submandibular gland
The gland is mixed serous and mucous although the serous element predominates (~90%). In mixed acini the serous cells form caps, or demilunes, on the periphery of the mucous cells. The intercalated ducts are shorter and the striated ducts more conspicuous than those of the parotid gland.

Sublingual gland
The gland is also mixed but is predominantly mucous in type. The mucous acini form elongated tubules with peripheral serous demilunes.

Minor salivary glands
These are most numerous at the junction of the hard and soft palate, lips and buccal mucosa. The minor glands of the lateral aspects of the tongue, lips and buccal mucosa are seromucous whereas those in the ventral tongue, palate, glossophyregeal area and retromolar pad are predominantly mucous. Salivary glands related to the circumvallate papillae (von Ebner’s glands) are serous in type. The minor glands are not encapsulated, and those in the tongue and lip especially can be deeply located in the musculature.

Epidemiology
The epidemiology of salivary gland tumours is not well documented (2053). In many studies the data are limited, as some are restricted to parotid gland neoplasms or tumours of major glands. In addition, most salivary gland tumours are benign and some cancer registries have only included malignant tumours. One study specifically excluded Warthin tumour, which is the second most common benign salivary neoplasm (698). In addition, several investigators felt that their quoted incidence figures were an underestimate, particularly for benign tumours (963,1471,2053). The global annual incidence when all salivary gland tumours were considered varied from 0.4-13.5 cases per 100,000 population (669). The frequency of malignant
salivary neoplasms ranged from 0.4-2.6 cases per 100,000 population {1353,1960,2053,2503}. In the United States, salivary gland malignancies accounted for 6% of head and neck cancers, and 0.3% of all malignancies {2167}. There is also some geographic variation in the frequency of tumour types. In studies of patients from Denmark and parts of Pennsylvania, about 30% of all parotid tumours were Warthin tumours, a sevenfold increase of the expected frequency {1765,2075}. The reported frequency of mucoepidermoid carcinomas among British patients (2.1%) is much lower than the worldwide range of 5-15% {703,704,1772,2580}. There was a very high reported incidence of salivary gland tumours in North American Inuits from 1950-1966 {1087,2255}. This was almost exclusively due to lymphoepithelial carcinomas that formed 25% of all malignancies in this population. Since then there has been a significant decline in the relative frequency of this tumour. A survey of different ethnic groups in Malaysia showed a higher frequency of salivary tumours in Malays than Chinese or Indians {1551}. Another study showed variations in the incidence of salivary tumours amongst different ethnic groups according to their city of residence {1705}. It should be noted that in some series malignant lymphoma and metastatic disease represent about 9% of major gland tumours, highlighting the need to include these neoplasms in differential diagnostic considerations {669,1916}.

**Site, age and sex distribution**

Between 64 and 80% of all primary epithelial salivary gland tumours occur in the parotid gland with most located in the superficial (lateral) lobe; 7-11% occur in the submandibular glands; fewer than 1% occur in the sublingual glands; and 9-23% occur in minor glands {669,679,703,2301,2439}. Benign tumours represent 54-79%, and 21-46% are malignant. The proportion of malignant tumours, however, varies greatly by site. Malignant tumours comprise 15-32% of parotid tumours, 41-45% of submandibular tumours, 70-90% of sublingual tumours, and 50% of minor gland tumours. Eighty to 90% of tumours that occur in the tongue, floor of mouth, and retromolar areas are malignant. Females are more frequently affected, but there is some gender variation according to the tumour type. The average ages of patients with benign and malignant tumours are 46 and 47 years, respectively, and the peak incidence of most of the specific types is in the sixth and seventh decades. However, the highest incidence of pleomorphic adenomas, mucoepidermoid carcinomas, and acinic cell carcinomas is in the third and fourth decades. In patients under 17 years of age, the frequency of mesenchymal tumours of the major glands is similar to that of epithelial tumours {1304,1413,2302,2337}. In this age group, pleomorphic adenomas, mucoepidermoid carcinomas and acinic cell carcinomas account for about 90% of epithelial tumours, and the frequency of benign and malignant tumours is essentially equal.

Among all patients, the most common tumour type is pleomorphic adenoma, which accounts for about 50% of all tumours. Warthin tumour is second in frequency among benign tumours and, in most large studies, mucoepidermoid carcinoma is the most common malignant tumour {669,679,703,2301,2439}. Most salivary neoplasms are classified into categories according to their histogenesis. A survey of surgical specimens from the parotid gland revealed that 54% of all parotid tumours were Warthin tumours, a sevenfold increase of the expected frequency {193,194,2543,2544}. The risk was directly related to the level of exposure to ionizing radiation. There was a high frequency of both mucoepidermoid carcinomas and Warthin tumours in these patients {2229}.

Therapeutic radiation, particularly of the head and neck region, has been linked with a significantly increased risk of developing salivary gland cancers {1725,1754,2197,2268}. There appears to be a risk from iodine131 used in the treatment of thyroid disease, as the isotope is also concentrated in the salivary glands {1111}. There is evidence that exposure to routine dental radiographs is associated with an increased risk of salivary gland carcinoma {2088,2089}. Exposure to ultraviolet radiation has also been implicated {1832,2451,2452}. There appears to be no excess risk in those exposed to radon {1733}, or the microwaves of cellular telephones {92,1224}.

**Occupation**

It has been shown that workers in a variety of industries have an increased incidence of salivary gland carcinomas. These include rubber manufacturing {1127,1620}, exposure to metal in the plumbing industry {1730} and nickel compounds {1127}, woodworking in the automobile industry {2512} and employment in hairdressing and beauty shops {2513,2514}. An increased risk of salivary gland cancers was reported in people living in certain Quebec counties where asbestos was mined, and the risk was inversely proportional to the distance from the mines {935}.

**Lifestyle and nutrition**

No association was found between tobacco use and alcohol consumption and salivary gland cancers in a case/control study {1801}, confirming previous findings {1295,2792}. One study showed an elevated risk in men but not women {1127}. However, there is a strong association between smoking and Warthin tumour (Section on Warthin tumour). Exposure to silica dust and kerosene as a cooking fluid increased...
the risk of developing salivary malignancy in a Chinese population [2902], and a higher level of risk of parotid carcinomas was associated with exposure to nickel, chromium, asbestos and cement dust in a European study [603]. An increased level of risk has been postulated in those with a high cholesterol intake [1128].

**Hormones**

Endogenous hormones have been reported in normal and neoplastic salivary glands, but some of the results have been conflicting. Estrogen receptors were found in nearly 80% of normal glands in males and females and four out of eight salivary tumours in women had estrogen receptor levels similar to those of "hornnormally dependent" breast carcinomas [606]. However, more recent studies have not confirmed this finding and questioned the methodology [616]. Estrogen receptors have been reported in a minority of cases of acinic cell carcinoma, mucoepidermoid carcinoma [1214] and salivary duct carcinoma [134], but were not detected in adenoid cystic carcinoma [616,1214,1732,2335]. Estrogen or estrogen receptors have been reported in pleomorphic adenomas in some studies [1214,1764,1946], but in others, estrogen receptors were absent [1851].

Progestrone receptors have been reported in normal salivary glands [892,2335]. They have been detected in a minority of pleomorphic adenomas [892,1214] but high levels of expression were reported in recurrent pleomorphic adenomas and this was thought to be a prognostic factor [892]. However, a recent study failed to show progesterone receptors in all the benign salivary tumours examined [1851]. Progesterone receptors were seen in 2/10 acinic cell carcinomas and 3/10 mucoepidermoid carcinomas [1214] but were not detected in salivary duct carcinoma [134]. They have been reported in adenoid cystic carcinomas in some studies [1214,1965] but in others they were absent, or present in only a few tumours [616,1214].

Androgen receptors are present in over 90% of salivary duct carcinomas [711,712,1265]. A recent study showed immunoreactivity for androgen receptors in all their cases of salivary duct carcinoma, carcinoma ex pleomorphic adenoma and basal cell adenocarcinoma [1851]. There was also staining for the receptors in a fifth of their cases of acinic cell carcinoma, mucoepidermoid carcinoma and adenoid cystic carcinoma.

**Diagnostic imaging**

Plain radiography and sialography are useful for ductal inflammatory disease, but computed tomography (CT), ultrasonography, CT sialography, and magnetic resonance imaging (MRI) are usually better for evaluation of suspected neoplastic disease. MRI is particularly useful when inflammatory disease is not suspected. It does not have the risks of radiation exposure nor complications with intraductal injection of contrast media, and it is often superior in demonstrating the interface of tumour and surrounding tissues. T1-weighted images of normal parotid have an image signal intermediate between fat and muscle whereas submandibular tissue is closer to muscle in intensity. With advanced age and fatty infiltration, the signal intensity of parotid tissue approaches fat. Most salivary gland tumours are brighter on T2 than T1 images but this difference is minimal in prominently cellular tumours. Lesions with higher water content, such as human immunodeficiency virus related parotid cysts, Warthin tumours, cystadenomas and cystadenocarcinomas, and cystic mucoepidermoid carcinomas, have a bright T2 signal.

**Fine needle aspiration biopsy**

Fine needle aspiration biopsy (FNA) can provide clinicians with rapid, non-surgical diagnoses. It can be performed at the time of initial consultation. Correlation of the clinical impression, cytologic diagnosis and radiographic imaging studies can then guide along different treatment pathways. FNA can be used both as a diagnostic test and as a screening tool to triage patients into different treatment groups i.e. surgical vs. medical management vs. to follow without intervention [2109]. FNA biopsy is useful in establishing whether a given lesion is inflammatory or neoplastic, is a lymphoma or an epithelial malignancy, or represents a metastasis or a primary tumour [424, 1585,2892]. Unnecessary surgery can be avoided in approximately one third of cases [668] especially in: (1) patients whose salivary gland lesion is part of a more generalized disease process, (2) inflammatory lesions where a clinical suspicion of malignancy is low, (3) patients in poor health who are not good operative candidates, (4) patients with metastasis to a salivary gland or adjacent lymph node, (5) some examples of lymphoproliferative disease [763] or (6) in a primary soft tissue or skin appendage lesion arising in the area of a major salivary gland. A number of series have examined the diagnostic accuracy of salivary FNA [26,495,2474,2887] with false positive and false negative rates ranging from 1-14%. The rate of correctly establishing a diagnosis as benign or malignant ranges from 81-98% in most recent reports. However, a specific diagnosis can only be made in approximately 60-75% of cases [668]. False negative diagnoses due to inadequate sampling appear to be the most frequent error.

**Frozen section examination**

When considering all head and neck sites, the accuracy of frozen section diagnoses of the salivary gland is the most controversial. A review of 2460 frozen sections from 24 series revealed an overall accuracy rate for a benign or malignant diagnosis, excluding deferred diagnoses, of 96% [379,900,1697,2170,2900]. False-positive rates (benign tumours initially diagnosed as malignant) were 1.1%, false-negative rates (malignant tumours initially diagnosed as benign) were 2.6%, and 2% of cases were deferred. If one subdivides the salivary gland lesions into benign and malignant groups, the accuracy rate (98.7%, excluding deferred diagnoses) is excellent for the benign lesions, which compose 80% of the frozen sections. However, in the malignant tumour group, the accuracy rate (85.9%) is suboptimal [900]. The most common benign tumour overdiagnosed as malignant was pleomorphic adenoma. This was frequently called mucoepidermoid carcinoma or adenoid cystic carcinoma [904]. Mucoepidermoid carcinoma is the malignancy most frequently associated with a false negative benign frozen section diagnosis, while acinic cell carcinoma, adenoid cystic carcinoma, carcinoma ex pleomorphic adenoma and an occasional lymphoma have also caused difficulty.

**Staging**

Staging of carcinomas of the major salivary glands is based on tumour size,
local extension of tumour, metastasis to regional nodes, and distant metastases (see TNM classification). Recent changes in the staging system include a revision in the definition of T3 and the division of T4 into tumours that are resectable (T4a) and unresectable (T4b) [947,2418]. According to TNM rules, tumours arising in minor salivary glands are classified according to the criteria for other carcinomas at their anatomic site of origin, e.g., oral cavity. Spiro and co-workers have successfully applied the criteria used for squamous cell carcinoma of the oral cavity, pharynx, larynx, and sinus to mucoepidermoid carcinoma [2305,2863].

Genetics

The goal of the molecular biological studies of salivary gland tumours is to define objective markers that may supplant the subjective phenotypic evaluation in the diagnosis, biological assessment and therapeutic stratification of patients with these tumours. The following molecular genetic events tentatively characterize some of these tumours:

1. Chromosomes 3p21, 8q12 and 12q13-15 rearrangements and the PLAG-1 and HMGI-C genes in pleomorphic adenomas
2. Translocations of chromosomes 11q21 and 19p13 in both Warthin tumour and mucoepidermoid carcinoma.
3. Structural and molecular alterations at 6q, 8q, 12q in adenoid cystic and carcinoma ex-pleomorphic adenoma.
4. Elevated HER-2 gene expression and gene amplification in mucoepidermoid, salivary duct and adenocarcinomas.

**EGFR**

Several studies have shown high expression of EGFR/HER-2/neu family members in mucoepidermoid and adenoid cystic carcinoma. The data suggest a biological role for members of this pathway in these tumours and their potential use as a target for therapy [887].

**C-erbB-2/HER-2/neu**

This is an oncogene that encodes for a transmembrane glycoprotein receptor involved in cell growth and differentiation. The gene is a member of the EGFR signal transduction family and has been shown to be overexpressed in aggressive breast cancer. Studies in salivary gland adenocarcinoma, including salivary duct and mucoepidermoid carcinoma, point to a general consensus on the association of HER-2 overexpression and adverse clinicopathologic features [725, 884,1058,2086,2198,2465].

**C-Kit**

This is a proto-oncogene that encodes a transmembrane receptor type tyrosine kinase that belongs to the colony-stimulating factor-1 (CSF-1) and platelet-derived growth factors (PDGF;4-6). Upon binding to its ligand, a signalling cascade is initiated to stimulate growth and differentiation of haematopoietic cells (835). Studies of C-kit in salivary gland tumours have largely focused on adenoid cystic carcinoma and findings vary considerably. C-kit expression appears to be restricted to adenoid cystic carcinoma [1215,2006] and myoepithelial carcinomas [1215] but absent in polymorphous low-grade adenocarcinoma [2006] and other types of salivary gland tumours [1215].

None of the highly expressed tumours manifested genetic mutations at exons 11 & 17. The results confirm a previous study and underscore that a mechanism for gene activation and other genetic alterations may play a role [1117]. A more recent study of this gene indicates high expression in other types of salivary gland neoplasms as well (adenoid cystic carcinoma, polymorphous low-grade adenocarcinoma and monomorphic types of adenoma) [636].

**TP53**

TP53 is a tumour suppressor gene located at the short arm of chromosome 17. The protein product acts as a transcription factor for cell differentiation, proliferation and death [636,1117,1485]. The role of this gene in salivary gland tumorigenesis remains unknown. Studies of different tumours have yielded variable results [554,1327,2198]. The incidence of p53 expression in other benign, malignant and hybrid tumours is low and does not correlate with recurrence [1823]. At present there is insufficient information on the correlation between p53 and outcome. These unsettling results reflect the lack of technical and interpretative uniformity in assessing this marker [2421].

Expression profiles

A study of nine benign (4 Warthin tumours and 5 pleomorphic adenomas) and three carcinomas (2 mucoepidermoid and one clear cell carcinoma), using cDNA of 19,000 human expressed sequence tags, identified a small set of genes that separate mucoepidermoid and clear cell carcinomas from normal and benign counterparts. Genes identified in carcinomas were apoptosis related [802].

A study of adenoid cystic carcinoma using oligonucleotide array platform for 8920 human genes was recently reported [818]. The study identified a set of genes that included basement membrane and extracellular matrix-related genes and genes encoding transcription factors SOX4 and AP-2a and members of the Wnt/β-catenin signalling pathway. A recent study of the gene expression in a cohort of pleomorphic adenomas and in spectrum of malignant tumours has also delineated a potential genetic profile that may be used in the biological investigation of these tumours [1652].

Genetic susceptibility

There is no evidence of familial clustering. An association has been reported with dermal cylindromatosis in the setting of Brooke-Spiegler syndrome [1248].

Prognosis and predictive factors

Prognosis correlates most strongly with clinical stage, emphasizing the importance of early diagnosis. The microscopically grade and tumour type have been shown to be independent predictors of behaviour and often play an important role in optimizing treatment [1264,1918,2440,2447,2449,2519]. Locoregional failure of some types of salivary carcinomas results in a greater likelihood of distant metastasis indicating a need for aggressive initial surgery [2441]. As might be expected, there is often a positive correlation between grade and clinical stage.
Acinic cell carcinoma

Definition
Acinic cell carcinoma is a malignant epithelial neoplasm of salivary glands in which at least some of the neoplastic cells demonstrate serous acinar cell differentiation, which is characterized by cytoplasmic zymogen secretory granules. Salivary ductal cells are also a component of this neoplasm.

ICD-O code 8550/3

Synonyms
Acinic cell adenocarcinoma, acinous cell carcinoma. Acinic cell tumour is an inappropriate synonym since the malignant biologic behaviour of this neoplasm is well-established [2304].

Epidemiology
Slightly more women than men are affected. There is no predilection for any ethnic group. Affected patients range from young children to elderly adults with a fairly even distribution of patients from the second to the seventh decades of life. Four percent of the patients are under 20 years old [668,1304,1954].

Localization
The overwhelming majority, almost 80%, of acinic cell carcinomas occur in the parotid gland, and about 17% involve the intraoral minor salivary glands. Only about 4% develop in the submandibular gland, and less than 1% arise in the sublingual gland [668,2711,2886].

Clinical features
They typically manifest as slowly enlarging, solitary, unfixed masses in the parotid region, but a few are multinodular and/or fixed to skin or muscle. A third of patients also experience pain, which is often vague and intermittent, and 5-10% develop some facial paralysis. While the duration of symptoms in most patients is less than a year, it can be up to several decades in some cases [478,668,670,1435,2445].

Macroscopy
Most are 1-3 cm in largest dimension. They are usually circumscribed, solitary nodules, but some are ill-defined with irregular peripheries and/or multinodularity. The cut surface appears lobular and tan to red. They vary from firm to soft and solid to cystic.

Tumour spread
Usually, acinic cell carcinomas initially metastasize to cervical lymph nodes and subsequently to more distant sites, most commonly the lung [670,960].

Histopathology
While serous acinar cell differentiation defines acinic cell carcinoma, several cell types and histomorphologic growth patterns are recognized. These are acinar, intercalated ductal, vacuolated, clear, and non-specific glandular and solid/lobular, microcystic, papillary-cystic, and follicular growth patterns [161, 478,668,1492,2290,2304]. Acinar cells are large, polygonal cells with lightly basophilic, granular cytoplasm and round, eccentric nuclei. The cytoplasmic zymogen-like granules are PAS positive, resistant to diastase digestion, and weakly stained or non-stained with mucicarmine. However, the PAS positivity can sometimes be very patchy and not immediately obvious. Intercalated duct type cells are smaller, eosinophilic to amphophilic, cuboidal with central nuclei, and surround variably sized luminal spaces. Vacuolated cells contain clear, cytoplasmic vacuoles that vary in number and size. The vacuoles are PAS negative. Clear cells are similar in size and shape to acinar cells but have non-staining cytoplasm that is non-reactive with PAS staining. Non-specific glandular cells are round to polygonal, amphophilic to eosinophilic cells with round nuclei and poorly demarcated cell borders. They often develop in syncytial sheets. Tumour cells are closely apposed to one another in sheets, nodules, or aggregates in the solid/lobular growth pattern. Numerous small spaces that vary from several microns to a millimetre or more in size characterize the microcystic pattern. Prominent cystic lumina, larger than the...
Acinic cell carcinoma

Microcystic spaces that are partially filled with papillary epithelial proliferations characterize the papillary-cystic pattern. This variant, in particular, may be very vascular and haemorrhagic and sometimes phagocytosis of haemosiderin by luminal tumour cells is a conspicuous feature. In the follicular pattern, multiple, epithelial-lined cystic spaces are filled with eosinophilic proteinaceous material, which produces a thyroid follicle-like appearance. Psammoma bodies are occasionally seen and are sometimes numerous. They are not restricted to the papillary-cystic variant and have been reported in FNA specimens. Although a single cell type and growth pattern often dominate, many tumours have combinations of cell types and growth patterns. Acinar cells and intercalated duct-like cells often dominate while the other cell types seldom do. Clear cells are seen in only 6% of all acinic cell carcinomas [670]. They are usually focal and only rarely cause diagnostic confusion. The solid/lobular and microcystic patterns are most frequent, followed by the papillary-cystic and follicular patterns.

A prominent lymphoid infiltrate of the stroma is associated with many acinic cell carcinomas [83,1717]. Whereas a heavy lymphoid infiltrate by itself has no prognostic significance, some tumours are well-circumscribed masses arranged in a microfollicular growth pattern and with a low proliferation index. They are completely surrounded by the lymphoid infiltrate (with germinal centre formation) and a thin fibrous pseudocapsule. These tumours appear to constitute a subgroup that behaves far less aggressively than other acinic cell carcinomas [1717].

Immunoprofile

Although the immunoprofile is non-specific, acinic cell carcinomas are reactive for cytokeratin, transferrin, lactoferrin, alpha 1-antitrypsin, alpha 1-antichymotrypsin, IgA, carinoembryonic antigen, Leu M1 antigen, cyclooxygenase-2, vasoactive intestinal polypeptide, and amylase. The zymogen granules in the neoplastic acinar cells are often non-reactive with anti-α-amylase immunostain, an enzyme in zymogen granules of normal serous acinar cells. Reactivity for oestrogen receptor, progesterone receptor, and prostate-specific antigen has been described in some tumours [338, 429,995,1031,1049,1214,2230,2296, 2529,2571]. Approximately 10% of tumours are positive for S-100 protein [2529].

Electron microscopy

Multiple, round, variably electron dense, cytoplasmic secretory granules characterize acinar type cells. The number and size of the granules varies. Rough endoplasmic reticulum, numerous mitochondria, and sparse microvilli are also typically. Some cells contain vacuoles of varying size and shape. Basal lamina separates groups of acinar and ductal

**Fig. 5.2** Acinic cell carcinoma. **A** Clear cells in acinic cell carcinoma are similar in size and shape to acinar-type cells but have non-staining cytoplasm. Some cells have a variable amount of eosinophilic cytoplasm. **B** The cytoplasmic granules in serous acinar-type cells in acinic cell carcinoma stain with PAS and are resistant to diastase digestion. **C** Sheets of tumour cells, acinar-type cells in this case, with few or no cystic spaces characterizes a solid growth pattern in acinic cell carcinoma. **D** Often acinar type cells are scattered among nonspecific glandular cells. They are often inconspicuous with H&E-stain but highlighted with PAS stain (PAS stain).
tumour cells from the stromal tissues. The light microscopically clear cells are the result of artefactual changes or dilatations of rough endoplasmic reticulum, lipid inclusions, enzymatic degradation of secretory granules, and intracytoplasmic pseudolumina [398,539,543,971].

**Histogenesis**
Most investigators consider that these tumours arise from neoplastic transformation of the terminal duct cells (intercalated duct cells) with histodifferentiation toward serous acinar cells. It has been shown [539,540], however, that normal serous acinar cells undergo mitotic division, and some acinic cell carcinomas could arise from transformation of these cells.

**Genetics**

**Cytogenetics**
Multiple structural and numerical abnormalities of these tumours have been reported but no consistent or specific alterations can be defined. Deletions of chromosome 6q, loss of Y and trisomy 21 have been reported [2238]. A recent report of multiple analyses from one tumour showed various structural abnormalities, suggesting a polyclonal derivation [1218].

**Molecular genetics**
In the largest molecular analysis of these tumours 21 (84%) of the 25 tumours studied showed LOH in at least one of the 20 loci on chromosomes 1,4,5,6 and 17 [647]. The most frequently altered regions were noted at chromosomes 4p, 5q, 6p and 17p regions. Chromosomes 4p15-16, 6p25-qter and 17p11 showed the highest incidence of alterations.

Another study of multiple spatially obtained samples from one tumour showed evidence for polyclonality suggesting different origins for this tumour [1218].

**Prognosis and predictive factors**
The average among several studies is a recurrence rate of about 35% and a metastatic rate and disease-associated death incidence of about 16% [441,478,670,960,1060,1112,1492,1845,1938,2017,2445]. Multiple recurrences and metastasis to cervical lymph nodes indicate a poor prognosis. Distant metastasis is associated with very poor survival. While tumours in the submandibular gland are more aggressive than those in the parotid gland, acinic cell carcinomas in minor salivary glands are less aggressive than those in the major salivary glands [340,864,1112,2886].

Attempts at histological grading have been controversial and inconsistent. Features that are often associated with more aggressive tumours include frequent mitoses, focal necrosis, neural invasion, pleomorphism, infiltration, and stromal hyalinisation [161,650,670,960,2017,2445]. Occasional cases of dedifferentiation from a low-grade to a high-grade malignancy have been reported. These tumours are characterized by cytological pleomorphism, increased mitotic and proliferation indices and have a worse prognosis [594,1063,1911,2459].

Staging is often a better predictor of outcome than histomorphologic grading. Large size, involvement of the deep lobe of the parotid gland, and incomplete resection indicate a poor prognosis. The cell proliferation marker Ki-67 has shown the most promise as a predictor of biological behaviour. No recurrences of acinic cell carcinomas were seen when the percentage of positively immunostained tumour cells was below 5% whereas most patients with tumour indices above 10% had unfavourable outcomes [1060,2388].

**Fig. 5.3** Acinic cell carcinoma. A Varibly sized luminal or cystic spaces, usually microscopic rather than macroscopic, identifies the microcystic pattern of acinic cell carcinoma. B Microcystic and macrocystic spaces, surrounded by tumour cells and filled with eosinophilic proteinaceous material, resemble thyroid follicles and characterize the follicular pattern, which is an infrequent variant of acinic cell carcinoma. C A prominent lymphocytic infiltrate of the stroma, usually with lymphoid follicles, is common and should not be misinterpreted as evidence of lymph node metastasis.

**Fig. 5.4** Acinic cell carcinoma. High-grade, poorly differentiated carcinoma (right) and typical acinic cell carcinoma (left) in a single neoplasm has been designated as “dedifferentiated” acinic cell carcinoma.
Mucoepidermoid carcinoma

Definition
Mucoepidermoid carcinoma is a malignant glandular epithelial neoplasm characterized by mucous, intermediate and epidermoid cells, with columnar, clear cell and oncocytoid features.

ICD code 8430/3

Synonyms
Mixed epidermoid and mucus secreting carcinoma. Mucoepidermoid tumour is an inappropriate synonym since the malignant biologic behaviour of this neoplasm is well established.

Epidemiology
Mucoepidermoid carcinoma (MEC) is the most common primary salivary gland malignancy in both adults and children (1560,2681,2711). MEC demonstrates a wide, nearly uniform age distribution, with diminution in paediatric and geriatric life (456,1850). Mean patient age is approximately 45 years. Sixty percent of palate lesions are in patients under 40. Tongue neoplasms are reported at an older average age. There is a 3:2 female predilection, but higher female predominance for tongue and retromolar pad tumours (668).

Localization
Approximately half of tumours (53%) occur in major glands. The parotid glands predominate, representing 45%, with 7% for submandibular glands and 1% in sublingual glands. The most frequent intra-oral sites are the palate and buccal mucosa.

Clinical features
Signs and symptoms
Most tumours present as firm, fixed and painless swellings. Sublingual gland lesions may demonstrate pain in spite of small size. Superficial intraoral neoplasms may exhibit a blue-red colour and mimic a mucocele or vascular lesion. The mucosa overlying palatal tumours can be papillary. Cortical bone is sometimes superficially eroded. Symptoms can include pain, otorrhoea, paraesthesia, facial nerve palsy, dysphagia, bleeding and trismus (703).

Macroscopy
Tumours are firm, smooth, often cystic, tan, white or pink with well-defined or infiltrative edges.

Tumour spread and staging
Parotid gland tumours spread to adjacent pre-auricular lymph nodes, then to the submandibular region. Submandibular gland neoplasms spread to submandibular and the upper jugular lymphatic chain. Palatal lesions may extend into the upper respiratory tract and skull base. Lip lesions invade submental nodes and intraoral tumours metastasize to submandibular, post auricular and upper accessory nodes in neck level II. With advancing disease, levels III, IV and V may become involved. Distant metastases may be widespread to lung, liver, bone, and brain.

Histopathology
Mucoepidermoid carcinoma is characterized by squamoid (epidermoid), mucus producing and cells of intermediate type. The proportion of different cell types and their architectural configuration (including cyst formation) varies in and between tumours.

Table 5.1 Histopathologic features, point values and point scores used in grading mucoepidermoid carcinoma

<table>
<thead>
<tr>
<th>Histopathologic feature</th>
<th>Point value</th>
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<tbody>
<tr>
<td>Cystic component &lt; 20%</td>
<td>2</td>
</tr>
<tr>
<td>Neural invasion</td>
<td>2</td>
</tr>
<tr>
<td>Necrosis</td>
<td>3</td>
</tr>
<tr>
<td>4 or more mitoses / 10 hpf</td>
<td>3</td>
</tr>
<tr>
<td>Anaplasia</td>
<td>4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Tumour Grade</th>
<th>Point Score</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td>0 - 4</td>
</tr>
<tr>
<td>Intermediate</td>
<td>5 - 6</td>
</tr>
<tr>
<td>High</td>
<td>7 or more</td>
</tr>
</tbody>
</table>

Fig. 5.5 Mucoepidermoid carcinoma. A Low-grade. B Intermediate grade.
They are usually multicystic with a solid component and sometimes the latter predominates. Some tumours have defined borders but infiltration of gland parenchyma is evident. Cystic spaces are lined by mucous cells with basaloid or cuboidal intermediate cells interspersed, and to a lesser degree, polygonal epidermoid cells, but keratinization is rare. Mucous cells are large, with pale cytoplasm and peripherally displaced nuclei. They typically constitute less than 10% of the tumour. Sialomucin content is demonstrated by muciarmine or Alcian blue staining. Intermediate cells usually predominate. Clear, columnar and/or oncocytic cell populations may be present and occasionally are prominent [985, 1198, 1996]. Clear cells demonstrate minimal sialomucin, but are diastase-sensitive periodic acid-Schiff positive, indicating glycogen content [666]. Focal sclerosis and/or mucus extravasation with inflammation is common. A sclerosing variant has been described [2657]. Neural invasion, necrosis, increased mitoses or cellular anaplasia are uncommon. At the tumour edge, a lymphocytic infiltrate with possible germinal centre formation can mimic nodal invasion [83].

**Grading**

Several systems have been proposed to grade this neoplasm, but none has been universally accepted [86, 258, 695, 1850, 2443]. However, one recent system using five histopathologic features has been shown to be reproducible in defining low, intermediate and high-grade tumours [86, 972, 1766]. In the submandibular gland low-grade tumours tend to behave more aggressively [921].

**Immunoprofile**

Squamous cells may be sparse in mucoepidermoid carcinoma and high molecular weight cytokeratins can help identify them.

**Differential diagnosis**

Differential diagnosis includes necrotizing sialometaplasia [263], inverted ductal papilloma, cystadenoma [2292], carcinomas composed of clear cells, adenosquamous carcinoma, squamous cell carcinoma and metastases.

**Genetics**

**Cytogenetics**

Several MECs have been reported to possess t(11:19) (q21:p13) translocation as the only abnormality (or with other structural and numerical alterations). This abnormality is also shared by acute leukaemia [655, 1130, 1904].

**Molecular genetics**

Molecular studies of these tumours are few and limited in number of cases. They show infrequent genetic loss at chromosomes 9p21, 8q, 5p, 16q and 12p [351, 1228, 2408]. Studies of the H-ras gene in these tumours have reported 18% mutations at codon 12 and/or 13 (one case) and no mutations at codon 61 [2858]. The mutations are mainly found in high-grade tumours [2859]. Recently molecular analysis of the t(11:19) (q21;p12) resulted in the identification of a fusion transcript resulting from the binding of exon-1 of a novel gene of unknown function, the mucoepidermoid carcinoma translocated gene-1 (MECT1), at 19p13 region with exons 2-5 of a novel member of the mastermind-like gene family (MAML2) at 11q21 region. This transcript activate the notch target genes.

**Prognosis and predictive factors**

Most patients have a favourable outcome. In one study, 8% of patients died of disease: 11% and 5% for major and minor gland tumours, respectively. Death correlated with high-grade histopathologic features in minor gland and parotid gland tumours, but not in patients with submandibular gland tumours [921]. Death resulted from unresetable locoregional tumour, distant metastases or complications of adjunctive therapy [2609]. The impact of grading on prognosis was described before and, additionally a MIB-1 index >10% correlates with high histopathologic grade, increased recurrence, metastasis and decreased patient survival [2387, 2905]. Currently, there are no prognostically useful genetic factors.
Adenoid cystic carcinoma

Definition
Adenoid cystic carcinoma is a basaloid tumour consisting of epithelial and myoepithelial cells in variable morphologic configurations, including tubular, cribriform and solid patterns. It has a relentless clinical course and usually a fatal outcome.

ICD code 8200/3

Epidemiology
Adenoid cystic carcinomas (AdCC) comprise approximately 10% of all epithelial salivary neoplasms and most frequently involve the parotid, submandibular and minor salivary glands. They comprise 30% of epithelial minor salivary gland tumours with the highest frequency in the palate, followed by the tongue, buccal mucosa, lip and floor of mouth. The tumour occurs in all age groups with a high frequency in middle-aged and older patients. There is no apparent sex predilection except for a high incidence in women with submandibular tumours [1663,1849,2016,2444].

Clinical features
The most common symptom is a slow growing mass followed by pain due to the propensity of these tumours for perineural invasion. Facial nerve paralysis may also occur [1849,2016,2444,2519].

Macroscopy
The carcinomas are solid, well-circumscribed but unencapsulated. They present as light-tan and firm masses of variable sizes. They are invariably infiltrative [161,1663,2439].

Histopathology
Tumours consist of two main cell types: ductal and modified myoepithelial cells that typically have hyperchromatic, angular nuclei and frequently clear cytoplasm. There are three defined patterns: tubular, cribriform and solid. In the tubular form, well-formed ducts and tubules with central lumina are lined by inner epithelial and outer myoepithelial cells. The cribriform pattern, the most frequent, is characterized by nests of cells with cylindromatous microcystic spaces. These are filled with hyaline or basophilic mucoid material. The solid or basaloid type is formed of sheets of uniform basaloid cells lacking tubular or microcystic formation. In the cribriform and solid variants small true ducts are invariably present but may not be immediately apparent. Each of these forms can be observed as the dominant component or more commonly as a part of a composite tumour [161,1663,1849,2016,2444,2519]. The stroma within the tumour is generally hyalinized and may manifest mucinous or myxoid features. In some tumours there is extensive stromal hyalinization with attenuation of the epithelial component. Perineural and to a lesser extent, intraneural invasion is a common and frequently conspicuous feature of AdCC. Tumours can extend along nerves for a considerable distance beyond the clinically apparent boundaries of the tumour. In addition, the tumour may invade bone extensively before there is

Table 5.2 Differential diagnosis of adenoid cystic carcinoma

<table>
<thead>
<tr>
<th>Tumour type</th>
<th>Pattern</th>
<th>Cellular features</th>
<th>Perineural invasion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basal cell adenoma</td>
<td>Syncytial/ non-invasive</td>
<td>Uniform Basaloid</td>
<td>No</td>
</tr>
<tr>
<td>Epithelial-myoepithelial carcinoma</td>
<td>Tubular/biphasic</td>
<td>Uniform, with clear outer cells</td>
<td>Rare</td>
</tr>
<tr>
<td>Basaloid squamous cell carcinoma</td>
<td>Syncytial</td>
<td>Marked pleomorphism focal keratinization</td>
<td>Rare</td>
</tr>
<tr>
<td>Basal cell adenocarcinoma</td>
<td>Syncytial/invasive</td>
<td>Mild pleomorphism/ invasive</td>
<td>Yes</td>
</tr>
<tr>
<td>AdCC Solid</td>
<td>Syncytial</td>
<td>Mild pleomorphism</td>
<td>Yes</td>
</tr>
<tr>
<td>AdCC Tubular/c cribriform</td>
<td>Ductal/ cylindromatous</td>
<td>Uniform biphasic</td>
<td>Yes</td>
</tr>
<tr>
<td>PLGA</td>
<td>Tubular papillary pattern variable</td>
<td>Mild pleomorphism</td>
<td>Yes</td>
</tr>
<tr>
<td>Cellular PA</td>
<td>Syncytial</td>
<td>Uniform</td>
<td>No</td>
</tr>
</tbody>
</table>

AdCC: Adenoid cystic carcinoma; PLGA: Polymorphous low-grade adenocarcinoma; PA: Pleomorphic adenoma

The cribriform pattern, the most frequent, is characterized by nests of cells with cylindromatous microcystic spaces. These are filled with hyaline or basophilic mucoid material. The solid or basaloid type is formed of sheets of uniform basaloid cells lacking tubular or microcystic formation. In the cribriform and solid variants small true ducts are invariably present but may not be immediately apparent. Each of these forms can be observed as the dominant component or more commonly as a part of a composite tumour [161,1663,1849,2016,2444,2519]. The stroma within the tumour is generally hyalinized and may manifest mucinous or myxoid features. In some tumours there is extensive stromal hyalinization with attenuation of the epithelial component. Perineural and to a lesser extent, intraneural invasion is a common and frequently conspicuous feature of AdCC. Tumours can extend along nerves for a considerable distance beyond the clinically apparent boundaries of the tumour. In addition, the tumour may invade bone extensively before there is

Fig. 5.7 Adenoid cystic carcinoma. Cribriform pattern with mucopolysaccharide filled spaces.
radiographical evidence of osseous destruction.
Adenoid cystic carcinoma occasionally occurs with other different neoplasms (hybrid tumours) [505,1823,2297,2416]. Pleomorphic carcinomas and sarcomatoid transformation of adenoid cystic carcinoma have been reported, mostly in recurrent and metastatic disease [397, 418].

Immunohistochemistry
In differentiating between polymorphous low-grade adenocarcinoma and adenoid cystic carcinoma, Ki-67 immunostaining may be helpful [2680]. DNA content, C-kit and E-cadherin have been found to be associated with the biological behaviour of these tumours [636, 637,1215,1577]. Ki-67 and p53 have also been studied in these tumours [2844], but no clear association with outcome have been reported. C-kit overexpression and its biological implication remains unknown. None of these markers, however, have been validated. Estrogen and progesterone receptor positivity has been reported in adenoid cystic carcinoma but the biological significance is currently unknown.

Differential diagnosis
Pleomorphic adenoma, polymorphous low-grade adenocarcinoma, epithelial-myoeptithelial carcinoma, basal cell adenoma or adenocarcinoma and basaloid squamous carcinomas are the major entities to be differentiated from adenoid cystic carcinoma.

Genetics
Cytogenetics
The most consistent, although not exclusive, reported alterations have been at chromosomes 6q, 9p and 17p12-13 regions. The t(6;9) (q21-24;p13-23) has been reported in several tumours and is considered to be a primary event in at least a subset of these tumours [657, 1220,1906,2238].

Molecular genetics
Frequent losses at 12q (33%) 6q23-qter, 13q21-q22 and 19q regions (40%) have been reported [657]. A study of the 9p21 regions and the p16 gene found only one tumour with LOH at this region and no mutations of the gene [351]. A recent genomic study identified new markers that may be helpful in future investigation of these tumours. Promoter methylation of the p16 was found in 20% of these tumours [1653]. Studies of other genes have been equally non-conclusive. Alterations of the p53 and Rb genes have been reported but no alterations in the K-ras have been found [2843].

Other studies have failed to confirm the value of grading [2439,2444] and underscored the significance of tumour size and clinical stage as the most consistent predictors of clinical outcome in patients with these tumours [2442,2449]. The 5-year survival rate is approximately 35% but the long-term survival is poorer. Eighty to 90% of patients die of disease in 10-15 years [993,2016]. The local recurrence rate ranges from 16-85% in several series of these tumours. Recurrence is a serious sign of incurability. Lymph node involvement is uncommon but has been reported to range from 5-25% and typically from tumours of the submandibular gland and is often due to contiguous spread rather than metastasis. The incidence of distant metastasis is estimated to range from 25-55%. The lung followed by bone, brain and liver are the common sites. Only 20% of patients with distant metastasis survive 5-years. The influence of perineural invasion on survival has been contradictory [860]. Wide local and radical surgical excisions with and without post-operative radiation is the treatment of choice [54,339, 1849,2439,2444,2519]. Radiation alone or with chemotherapy in the treatment of recurrent or metastatic disease has shown limited success. Radiotherapy, however, has been shown to improve local control in cases with microscopic residual disease [2670]. The value of chemotherapy in these tumours is limited and remains to be proven.

![Fig. 5.8 Adenoid cystic carcinoma. A Tubular form, composed of inner epithelial ductal and outer myoepithelial cells. B Solid form. Tumour cells are small and basaloïd with scanty cytoplasm.](image-url)
Polymorphous low-grade adenocarcinoma

Definition
A malignant epithelial tumour characterized by cytologic uniformity, morphologic diversity, an infiltrative growth pattern, and low metastatic potential.

ICD-O code 8525/3

Synonyms
Terminal duct carcinoma, lobular carcinoma

Epidemiology
PLGA is the second most common intraoral malignant salivary gland tumour, accounting for 26% of all carcinomas [2711]. The female-to-male ratio is about 2:1. Patient age ranges from 16-94 years mean 59 years. Over 70% of the patients are between the ages of 50 and 70 years [342,697]. To date, only two tumours have been reported in the pediatric population [2641].

Localization
Approximately 60% of the cases have involved the palate. Other intraoral locations are the buccal mucosa, retromolar region, upper lip, and the base of the tongue [342,697]. Uncommon locations include major salivary and lacrimal glands, nasopharynx and nasal cavity [1299,2763].

Clinical features
A painless mass in the palate is the most common clinical sign. The duration of the lesion has varied from weeks to as much as 40 years [342]. Bleeding, telangiectasia, or ulceration of the overlying mucosa occurs occasionally.

Macroscopy
PLGA usually appears as a firm, circumscribed, but non-encapsulated, yellow-tan lobulated nodule up to several centimetres in greatest dimension (average 2.2 cm) [342].

Histopathology
PLGA is characterized by cytologic uniformity, histologic diversity, and an infiltrative growth pattern. The tumour cells are small to medium size and uniform in shape with bland, minimally hyperchromatic, oval nuclei and only occasional nucleoli. Mitoses are uncommon and necrosis is not typical. The striking feature of these carcinomas is the variety of morphologic configurations between tumours and within an individual tumour. The main microscopic patterns are: 1) lobular, 2) papillary or papillary-cystic (typically focal), 3) cribriform areas sometimes resembling those in adenoid cystic carcinoma, and 4) trabecular or small, duct-like structures lined by a single layer of cuboidal cells. The cells form concentric whorls or targetoid arrangements around blood vessels or nerves. Foci of oncocytic, clear, squamous or mucous cells may be found. Stroma may show areas of mucinosis or hyalinization. Despite the innocuous cytologic appearance, the neoplasm always invades adjacent soft tissues and is uncapsulated. Neurotropism is common in PLGA. Invasion of adjacent bone may be seen in tumours of the palate or mandible.

Immunohistochemistry
The neoplastic cells of PLGA are immunoreactive with antibodies to cytokeratin (100%), vimentin (100%), S-100 protein (97%), carcinoembryonic antigen (54%), glial fibrillary acidic protein (GFAP) (15%), muscle specific actin (13%), and epithelial membrane antigen (12%) [342,2011,2763]. Expression of galectin 3 has been reported to be significant in PLGA (2006). Bcl-2 is over expressed in most cases of PLGA [342,2011].

Differential diagnosis
The differential diagnosis includes pleomorphic adenoma (PA) and adenoid cystic carcinoma (AdCC), especially in small biopsy specimens. Unlike PLGA, PA is nearly always circumscribed and is composed of proliferating stromal, epithelial, and myoepithelial cells. It lacks the infiltrative, noncircumscribed character of PLGA. Although myxoid tissue is present in both tumours, the myxochondroid and chondroid areas present in PA are not evident in PLGA. Also, the typical benign plasmacytoid myoepithelial cells characteristic of palatal PA are seldom observed in PLGA. Staining with GFAP may be helpful in differentiating PA from PLGA.

The distinction between PLGA and AdCC is based primarily on cytologic features. Cells in PLGA are cuboidal or columnar. They have vesicular nuclei and often conspicuous eosinophilic cytoplasm without the basaloid features characteristic of AdCC. Papillary and fascicular growth patterns are extremely rare in AdCC. Furthermore, PLGA does not have large cribriform pseudocystic spaces that contain pools of haemosiderin. The solid cellular areas of PLGA lack nuclear pleomorphism, necrosis, increased mitotic activity, and the numerous tubular structures characteristic of the solid variant of AdCC. The potential discriminating value of immunohistochemistry between cases of PLGA and AdCC remains controversial [547], although some subtle differences may be apparent when series of these two neoplasms are studied [342,2006,2011,2391]. Proliferative cell marker rates in PLGA are usually less than 6.4% (mean values 1.6% and 2.4%) [2391]. However, a higher proliferative rate (average 7%) has been reported by others investigators [2011].

Genetics
Cytogenetic studies of this tumour are few. A total of 7 cases of which two were carcinoma ex-pleomorphic adenoma, have been reported. Alterations at 8q12 were found in two, 12q rearrangements in five, two showed a clonal t(6:9) (p21;p22) and one a monosomy 22 (1651). Cytogenetic alterations in PLGA have frequently displayed chromosome 12 abnormalities affecting the q arm and the p arm (1651).
**Prognosis and predictive factors**

The overall survival rate of patients with PLGA is excellent [164,342,696, 697,808]. A review of series with large numbers of cases and with long-term follow-up revealed a local recurrence rate between 9% and 17% and a regional metastases rate from 9-15% [342,697]. Distant metastases have seldom been reported [342,697]. Deaths attributed to tumour are unusual, and they occurred after prolonged periods [342,697]. In studies which accepted tumours with a predominant papillary configuration a higher incidence of cervical lymph node metastasis was reported [697]. The status of such tumours within the spectrum of PLGA is controversial. Dedifferentiation of PLGA has been reported and carries a less favourable prognosis. Such tumours should not be included under the rubric of typical PLGA [2368].

Treatment consists of complete surgical excision. Neck dissection should be added for those patients with cervical adenopathy.

**Cribriform adenocarcinoma of the tongue**

A possible variant is cribriform adenocarcinoma of the tongue, but it is not yet clear whether this represents a genuine entity or just an unusual growth pattern in PLGA, with which there appears to be some overlap [1718].

So far described only in one series, all cases presented with a mass in the tongue, usually the posterior part, and synchronous metastases in lateral neck lymph nodes, but no distant spread. There was an equal sex incidence and the mean age at presentation was 50.4 years (range 25-70).

The tumour grows beneath the surface epithelium and infiltrates soft tissue. It is divided by fibrous septa into lobules, which are solid or cribriform. A characteristic feature is that some nearly solid islands have a glomeruloid arrangement of broad microfollicular papillae separated from a layer of peripheral columnar cells by a narrow cleft. Small numbers of tubules are seen, and occasional spindling of tumour cells may occur. The nuclei are uniform, pale and often overlap, closely mimicking those of papillary carcinoma of the thyroid. Mitotic figures are sparse. No necrosis or significant haemorrhage is seen, and the stroma includes hyalinized areas, and rarely psammoma bodies. The tumours are positive for cytokeratin, and more patchily for S-100 protein. Myoepithelial markers, such as actin are either negative or only focally positive. Thyroglobulin staining is consistently negative.
EPITHELIAL-MYOEPITHELIAL CARCINOMA

Definition
A malignant tumour composed of variable proportions of two cell types, which typically form duct-like structures. The biphasic morphology is represented by an inner layer of duct lining, epithelial-type cells and an outer layer of clear, myoepithelial-type cells.

ICD-O code 8562/3

Synonyms
Adenomyoepithelioma [176], clear cell adenoma [494, 2228], glycogen-rich adenoma [913], glycogen-rich adenocarcinoma [1758], clear cell carcinoma [407].

Epidemiology
Epithelial-myoepithelial carcinoma (EMC) represents around 1% of the salivary gland tumours. It is more prevalent in women (F: M=2:1). The patients range in age from 13 to 89 years, with the peak incidence in the 6th and 7th decades [436, 493, 614, 784, 1580]. Only two cases have been reported in the paediatric group [436, 1775].

Localization
EMC occurs mostly in major salivary glands, mainly in the parotid (60%), but also in the minor glands of oral mucosa and the upper [436, 493, 614, 784, 1580] and lower respiratory tract [610, 1126, 1174, 2002].

Clinical features
EMC forms a painless, slow-growing mass. Tumours arising in minor glands frequently present as ulcerated, submucosal nodules and have less well-defined margins. Rapid growth, facial nerve palsy and/or associated pain are suggestive of concomitant high-grade areas.

Macroscopy
EMC is characteristically a multinodular mass, with expansive borders and lacking a true capsule. Cystic spaces may be present. Tumours of the minor glands are poorly circumscribed.

Histopathology
EMC has a lobulated growth pattern with a mixed tubular and solid architectural arrangement. Papillary and cystic areas can be identified in around 20% of the cases. Tumours from minor salivary and sero-mucinous glands show infiltration of surrounding tissues and there is ulceration of the overlying mucosa in about 40% of the cases. The hallmark of EMC histology is the presence of bi-layered duct-like structures: the inner layer is formed by a single row of cuboidal cells, with dense, finely granular cytoplasm and central or basal, round nucleus. The outer layer may show single or multiple layers of polygonal cells, with well-defined borders; the cytoplasm is characteristically clear and the nucleus is vesicular and slightly eccentric. The double-layered pattern is preserved in papillary-cystic areas but solid tumour areas may be exclusively formed by clear cells. PAS positive, hyaline, eosinophilic strands of basement membrane-like material surround the duct-like structures and, in solid areas, divide the clear cells into theques. Coagulative necrosis at the centre of tumour nodules is uncommon. In rare cases, squamous differentiation and spindle cells are observed as well as an oncocytic appearance in the inner cell layer of neoplastic ducts. Perineural and vascular invasion are frequent and bone invasion may occur. None to 1-2 mitoses per 10 HPF can be identified in the clear cell population of EMC. Rare cases of dedifferentiation have been reported [42, 783].

Immunoprofile
Myoepithelial markers (smooth muscle actin, HHF35, p63 and/or calponin) stain the clear cell compartment. The luminal cells stain with cytokeratins.

Fig. 5.10  Epithelial-myoepithelial carcinoma (EMC) of the parotid gland. A Double layered architecture formed by an inner layer of eosinophilic cuboidal cells and an outer layer of clear, myoepithelial-type cells. B Dedifferentiated EMC. There is co-existence of areas typical of EMC (left) with areas of undifferentiated carcinoma (right).
Differential diagnosis
The differential diagnosis of EMC includes all primary salivary gland tumours that are predominantly formed by clear cells: pleomorphic adenoma, myoepithelioma, oncocytoma and mucoepidermoid carcinoma. Differential diagnosis with clear cell carcinoma, NOS relies on the demonstration of the peculiar amyloid-like quality of the stroma and on the absence of myoepithelial markers. Metastatic kidney and thyroid carcinoma may be distinguished using immunohistochemistry; CD10 and high-molecular weight cytokeratin in the former and thyroglobulin in the latter. EMC foci can be encountered within carcinoma ex pleomorphic adenoma as part of the carcinomatous component.

Genetics
A limited number of cases (6) have been karyotyped [656,1650,1751], half of them showing non-distinctive chromosomal alterations and the remaining normal karyotypes.

Prognosis and predictive factors
Recurrence occurs in around 40% of cases and metastasis in 14%. The most common metastatic sites are cervical lymph nodes, lung, liver and kidney. Death from disease complications occurs in less than 10% of the patients [436,493,614,784,1580]. Five- and 10 year overall survival rates are 80% and 72%, respectively [784]. Size and rapid tumour growth are associated with worse prognosis [42,783]. Margin status is a major pathological prognostic factor. Incomplete surgical excision is associated with recurrence and metastasis. The poorer prognosis associated with tumours located in minor salivary glands may be due to the higher frequency of recurrences due to incomplete surgery. Atypia is associated with unfavourable outcome [784] whenever present in more than 20% of tumour area. EMC is usually diploid [784,992]. Aneuploidy and high mitotic counts have been reported in cases with unfavourable prognosis [784]. Areas of dedifferentiation also predict poor outcome, with recurrence and metastasis in 70% of patients [42,783].
Clear cell carcinoma, not otherwise specified

**Definition**
Clear cell carcinoma, not otherwise specified (NOS), is a malignant epithelial neoplasm composed of a monomorphic population of cells that have optically clear cytoplasm with standard haematoxylin and eosin stains. Because many types of salivary gland neoplasms commonly or consistently have a component of clear cells, clear cell carcinoma is distinguished by the absence of features characteristic of these other neoplasms and its monomorphous population of clear cells.

**ICD-O code** 8310/3

**Synonyms**
Clear cell adenocarcinoma; hyalinizing clear cell carcinoma.
Clear cell carcinoma has been confused with epithelial-myoepithelial carcinoma (EMC), and EMC have been reported as clear cell carcinoma [407].

**Epidemiology**
The peak occurrence is in patients in the 40-70 year age range, and they are rare in children [668,2658]. There is no sex predilection.

**Localization**
Clear cell carcinomas are more frequent in the intraoral minor salivary glands than the major salivary glands [668,1728, 1931,2179,2369,2716]. The palate is most frequently involved, but buccal mucosa, tongue, floor of the mouth, lip, and retromolar and tonsillar areas are also affected.

**Clinical features**
The only sign in most cases is swelling, but mucosal ulceration and pain occur with some tumours. Patients have reported the durations of their tumours as 1 month to 15 years [2369].

**Macroscopy**
Although the size of the primary tumour is usually 3.0 cm or less, the tumours usually are poorly circumscribed and infiltrate adjacent salivary gland, mucosa, soft tissues, bone, and nerves. The cut surface is greyish-white.

**Histopathology**
A monomorphic population of polygonal to round cells with clear cytoplasm characterizes clear cell carcinomas. In some cases, a minority of cells have pale eosinophilic cytoplasm. Nuclei are eccentric and round and frequently contain small nucleoli. PAS staining with and without prior diastase digestion of the tissue demonstrates cytoplasmic glycogen that varies from marked to not evident. The adjective glycogen-rich has been used by some to identify clear cell carcinomas with a prominent glycogen content [1028,2658]. With mucicarmine stain, intracytoplasmic mucins are usually absent. The tumour cells are arranged in sheets, nests, or cords, and ductal structures are absent. Mitotic figures are rare, but some tumours have a moderate degree of nuclear pleomorphism. In the hyalinizing type, the stroma is composed of thick bands of hyalinized collagen [727,1728], but in other tumours it consists of interconnecting, thin fibrous septa that may be cellular or loosely collagenous. Clear cell carcinomas are unencapsulated and infiltrative.

**Immunoprofile**
While tumours are immunoreactive for cytokeratin, at least focally, immunohistochemical studies have given variable results for S100 protein, glial fibrillary acidic protein, actin, and vimentin [1028, 1728,1931,2348,2369,2658,2716]. Tumours that demonstrate histologic and immunohistochemical features of myoepithelial differentiation are best classified as clear cell variants of myoepithelioma or myoepithelial carcinoma [1719].

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**Fig. 5.13** Clear cell carcinoma. A Polygonal cells with nonstaining cytoplasm and absence of ductal lumens characterize clear cell carcinoma. B This example is composed of a mostly solid sheet of tumour cells with little stroma.
Electron microscopy
Tight junctions, desmosomal attachments, tonofilaments, microvilli, and basal lamina are features of duct cell differentiation [399,1028,1728,1758,1931,2369,2716].

Histogenesis
Ultrastructural investigations have found features of ductal but not myoepithelial differentiation.

Prognosis and predictive factors
Prognosis is excellent. A few tumours have metastasized to cervical lymph nodes and, rarely, the lung, but no patients have succumbed to this neoplasm [155,948,1728,2716].
Basal cell adenocarcinoma

Definition
Dominated by basaloid epithelial cells, basal cell adenocarcinoma is cytologically and histomorphologically similar to basal cell adenoma but is an infiltrative epithelial neoplasm with potential for metastasis.

ICD-O code 8147/3

Synonyms
Basaloid salivary carcinoma, carcinoma ex monomorphic adenoma, malignant basal cell adenoma, malignant basal cell tumour, and basal cell carcinoma [159, 408, 698, 1163, 1340, 1576]. Tumours in infants reported as basal cell adenoma/carcinoma or hybrids are best classified as sialoblastomas.

Epidemiology
There is no sex predilection. The average age of patients is 60 years, and only adults have been affected [668, 673, 1576, 1792, 2726].

Localization
Over 90%, of these tumours occur in the parotid gland, and they are rare in the minor salivary glands of the oral cavity [668, 673, 2703].

Clinical features
Rarely, patients complain of pain or tenderness; most tumours are asymptomatic except for swelling. The duration of tumours before excision ranges from weeks to years. Similar to some patients with basal cell adenomas, patients with basal cell adenocarcinomas may have a diathesis of multiple skin adnexal tumours and parotid basal cell adenocarcinomas [65, 668, 673, 1163, 1576].

Macroscopy
Basal cell adenocarcinomas most frequently occur in the superficial (lateral) lobe of the parotid gland. The cut surface has variable coloration of grey, tan-white, or brownish. The texture is homogeneous although some tumours are focally cystic. They are unencapsulated, but some tumours appear well-circumscribed while others are obviously infiltrative.

Histopathology
Basaloid epithelial cells, which vary from small, dark cells to larger, paler stained cells, form histomorphologic patterns that are described as solid, membranous, trabecular and tubular. A solid pattern, in which variable sized and shaped nests are separated by thin septa or thick bands of collagenous stroma, is most frequent. In the membranous type, tumours produce excessive amounts of eosinophilic, hyalinized basal lamina material that forms intercellular droplets and peripheral membranes. Interconnecting bands of basaloid cells characterize the trabecular growth pattern. In the tubular type, there are luminal spaces among the basaloid cells. There are foci of squamous differentiation in some tumours. The nuclei of tumour cells along the interface with the collagenous stroma are often palisaded. The degree of cytologic atypia and the number of mitotic figures varies from one tumour to another but is often quite minimal. Infiltration of tumour cells into parotid parenchyma, dermis, skeletal muscle, or periglandular fat distinguishes basal cell adenocarcinoma from basal cell adenoma. Vessel or peripheral nerve invasion is evident in about a fourth of the tumours.

Immunoprofile
Immunohistochemical staining is variable among tumours. Tumour cells are reactive for cytokeratins and often focally reactive for S100 protein, epithelial membrane antigen, and carcinoembryonic antigen. Limited reactivity for smooth muscle actin and vimentin supports myoepithelial differentiation of some cells [2097, 2793].

Fig. 5.15 Basal cell adenocarcinoma. A Basal cell adenocarcinoma, parotid gland. Invasive growth. B Abundant, prominently eosinophilic basal lamina material within and around nests of tumour is characteristic of the membranous pattern of basal cell adenocarcinoma.
Precursor lesions
Most basal cell adenocarcinomas probably develop de novo, but some arise by malignant transformation in basal cell adenomas [1576,1792].

Genetics
Cytogenetics
Chromosomal gains at 9p21.1-pter, 18q21.1-q22.3, and 22q11.23-q13.1 as well as losses at 2q24.2 and 4q25-q27 have been described [2612]. The gain at 22q12.3-q13.1 is described as also common in adenoid cystic carcinoma.

Molecular genetics
A study of two familial cases and two sporadic basaloid tumours for alterations at the 16q12-13 regions showed high frequency (80%) of LOH in both sporadic and familial basaloid tumours and dermal cylindromas of the familial cases. The minimally deleted region contained the CYLD gene. This study indicates that these tumours share the same alterations as dermal cylindromas and implicates the CYLD gene in their development [437].

Prognosis and predictive factors
While they are locally destructive and often recur, basal cell adenocarcinomas only occasionally metastasize, and death of patients is rare [408,673,698,1163,1340,1576,1792,1799]. Ki-67 and PCNA indices are low [782,2097].
Malignant sebaceous tumours

Sebaceous carcinoma

Definition
Sebaceous carcinoma is a malignant tumour composed of sebaceous cells of varying maturity that are arranged in sheets and/or nests with different degrees of pleomorphism, nuclear atypia and invasiveness.

ICD-O code 8410/3

Epidemiology
There is a bimodal age distribution with a peak incidence in the third decade and the 7th and 8th decades of life (range 17-93 years) \{669,896,901\}. The male and female incidence is almost equal. Unlike sebaceous neoplasms of the skin \{1132,2214\}, there is no increased risk of developing a visceral carcinoma in patients with a salivary gland sebaceous tumour.

Localization
Approximately 90% arise in the parotid area, with occasional tumours in the oral cavity, vallecula, sublingual gland, submandibular gland and epiglottis \{107,602,669,693,896\}.

Clinical features
Patients typically present with a painful mass with varying degrees of facial nerve paralysis and occasional fixation to the skin.

Macroscopy
Tumours have ranged from 0.6-8.5 cm in greatest dimension and vary from yellow, tan-white, greyish-white, white, to pale pink {896}. They are well circumscribed or partially encapsulated, with pushing or locally infiltrating margins.

Histopathology
Tumours are composed of multiple large foci and nests of cells with hyperchromatic nuclei and abundant clear to eosinophilic cytoplasm. Cellular pleomorphism and cytologic atypia are present to varying degrees and are much more prevalent than in sebaceous adenomas. Squamous differentiation is common. There may be areas of basaloId differentiation, particularly at the periphery of cellular nests. Areas of necrosis and fibrosis are common. Perineural invasion is seen in greater than 20% of tumours; vascular invasion is infrequent. Rare oncocyes and foreign body giant cells with histiocytes may be observed, but lymphoid tissue with follicles or subcapsular sinuses is not seen.

Prognosis and predictive factors
The treatment of choice is wide surgical excision for low stage carcinomas. Adjunctive radiation therapy is recommended for higher-stage and grade tumours. The overall 5-year survival rate is 62% {669,896}, slightly less than the survival for similar tumours arising in the skin and orbit (84.5%) {234}.

Sebaceous lymphadenocarcinoma

Definition
Sebaceous lymphadenocarcinoma is the malignant counterpart of sebaceous lymphadenoma. It is a carcinoma arising in a sebaceous lymphadenoma.

ICD-O code 8410/3
Synonym Carcinoma ex sebaceous lymphadenoma.

Epidemiology
It is the rarest salivary gland sebaceous tumour. To date, only three have been reported {901,1525}. All three patients were in their seventh decade; two patients were male and one female.

Localization
The tumours arose within the parotid gland or in periparotid lymph nodes.

Clinical features
Patients had histories of a mass, two of which were present for more than 20 years.

Macroscopy
Tumour colour varies from yellow-tan to grey.

Histopathology
These carcinomas are partially encapsulated and locally invasive with foci of sebaceous lymphadenoma intermixed with or adjacent to regions of pleomorphic carcinoma cells exhibiting varying degrees of invasiveness. The malignant portion has ranged from sebaceous carcinoma to sheets of poorly differentiated carcinoma, with areas of ductal differentiation, adenoid cystic carcinoma-like areas or foci of epithelial-myoepithelial carcinoma. Perineural invasion, collections of histiocytes and a foreign body giant cell reaction may occur. Cellular atypia is not observed in the sebaceous lymphadenoma portion of the tumour.

Fig. 5.18 A Sebaceous carcinoma. Solid growth of pleomorphic sebaceous tumour cells. Inset: Tumour cells are positive for fat stain (Sudan-III). B Sebaceous lymphadenocarcinoma composed of poorly differentiated carcinoma cells with areas of ductal differentiation.
**Definition**
Cystadenocarcinoma is a rare malignant tumour characterized by predominantly cystic growth that often exhibits intraluminal papillary growth. It lacks any additional specific histopathologic features that characterize the other types of salivary carcinomas showing cystic growth. It is conceptually the malignant counterpart of the benign cystadenoma.

**ICD-O code**
8440/3

**Synonyms**
Papillary cystadenocarcinoma, mucus-producing adenopapillary (non-epidermoid) carcinoma {224,679,2463}, malignant papillary cystadenoma {2133}, and low-grade papillary adenocarcinoma of the palate {38,1742,2784}.

**Epidemiology**
There is no sex predilection. The average age of patients is 59 years; more than 70% are over 50 years of age {790}.

**Localization**
About 65% occur in the major salivary glands and most of these arise in the parotid. Involvement of the sublingual gland is proportionately greater than of other benign or malignant tumours {790}. The buccal mucosa, lips, and palate are the most frequently involved minor gland sites.

**Clinical features**
Cystadenocarcinomas usually manifest as a slowly growing, compressible asymptomatic mass. Tumours of the palate may erode bone.

**Macroscopy**
The tumours have multiple cystic spaces that are variable in size and often filled with mucin. They are grossly at least partially circumscribed and have ranged in size from 0.4-6 cm.

**Histopathology**
The tumours are usually well circumscribed but not encapsulated. Numerous haphazardly arranged cysts are evident that are partially filled with mucin, vary in shape and size, and have limited intervening fibrous connective tissue. Small solid neoplastic islands or duct-like structures may occur between the cysts or at the advancing front of the tumour. In about 75% of the cases the lumens of the cysts exhibit varying degrees of papillary proliferation. In either case, cell types that comprise the lining epithelium include, most often, small and large cuboidal, and columnar cells, but mucous, clear and oncocytic cells are occasionally noted. The columnar-rich tumours often predominate in the intraluminal papillary areas and account for their “gastrointestinal” appearance, but the cells usually fail to stain for neutral mucin. Although nucleoli are evident, the nuclei typically are uniformly bland and mitoses rare. However, a prerequisite for the diagnosis is that the cysts and smaller duct-like structures at least focally infiltrate the salivary parenchyma and surrounding connective tissue. The presence of ruptured cysts with haemorrhage and granulation tissue is common.

**Differential diagnosis**
Distinction from cystadenoma may be difficult and relies largely on identification of infiltrative growth into salivary parenchyma or surrounding tissues. Review of multiple sections is often helpful. Low-grade mucoepidermoid carcinoma is typically cystic but, unlike cystadenocarcinoma, usually has a wide variety of cell types and areas that are more solid than cystic. The papillary cystic variant of acinic cell carcinoma has focal acinar differentiation and a greater degree of epithelial proliferation. Epidermoid differentiation in cystadenocarcinomas is rare.

**Prognosis and predictive factors**
Cystadenocarcinoma is a low-grade adenocarcinoma treated by superficial parotidectomy, glandectomy of submandibular and sublingual tumours, and wide excision of minor gland tumours. Bone resection is performed only when it is directly involved by tumour {411,535,790,2350}. In a study of 40 patients with follow-up data, all were alive or had died of other causes, four suffered metastasis to regional lymph nodes, one at the time of diagnosis and one after 55 months, and three experienced a recurrence at a mean interval of 76 months {790}.

**Fig. 5.19** Cystadenocarcinoma. A Focal collections of lymphoid tissue are present. No significant papillary luminal growth. B Cystic spaces are lined by morphologically bland low cuboidal epithelium, and are separated by loosely arranged fibrous stroma. C Papillary cystadenocarcinoma.
Definition
A rare, cystic, proliferative carcinoma that resembles the spectrum of breast lesions from atypical ductal hyperplasia to micropapillary and cribriform low-grade ductal carcinoma in-situ.

Synonym
Low-grade salivary duct carcinoma

Epidemiology
To date, all but one tumour have been diagnosed in the parotid gland and one in the palate [259,578,899,2562]. There is a female predominance of 2:1.

Clinical features
Patients are usually elderly and all but one patient presented with cystic parotid tumours.

Histopathology
Low-grade cribriform cystadenocarcinomas (LGCCC) are unencapsulated, consisting of single or multiple cysts, accompanied by adjacent intraductal proliferation. The cysts are lined by small, multilayered, proliferating, bland ductal cells with finely dispersed chromatin and small nucleoli. Within the cystic areas, they typically are arranged in a cribriform pattern and frequently have anastomosing, intracyctic micropapillae lining the cavity, which may contain fibrovascular cores. Separate, smaller ductal structures are variably filled by proliferating ductal epithelium with cribriform, micropapillary and solid areas. The overall appearance is very similar to breast atypical ductal hyperplasia and low-grade ductal carcinoma in-situ. Many superficial cells contain cytoplasmic apocrine-type microvacuoles (PAS-positive/diastase-resistant) and/or fine yellow to brown pigment resembling lipofuscin. Focal invasion into the surrounding tissue can be seen, characterized by small solid islands and reactive inflammation and desmoplasia. Perineural or vascular invasion typically is not present. Cellular pleomorphism and mitotic figures are usually absent and necrosis is extremely uncommon. Occasional tumours may demonstrate transition from low to intermediate or high-grade cytology, with scattered mitotic figures and focal necrosis.

Immunoprofile
These tumours demonstrate strong, diffuse S100 positivity. Myoepithelial markers (calponin or smooth muscle actin) highlight cells rimming the cystic spaces, confirming the intraductal nature of most, or all, of each tumour. No myoepithelial cells are admixed within the proliferative cellular component. Those tumours studied for HER2-neu antigen are uniformly negative.

Variants
Originally, this tumour was reported as a low-grade variant of salivary duct carcinoma. However, as no data have accumulated definitely relating this entity to ductal carcinoma and since there frequently is a prominent cystic component, for the purposes of this WHO classification, the tumour is listed as a variant of cystadenocarcinoma.

Differential diagnosis
The following tumours require exclusion: papillary cystic variant of acinic cell carcinoma, (PCVACC) and other variants of cystadenocarcinoma. PCVACC contains vacuolated cells similar to the microvacuolated cells of LGCCC. However, the vacuoles of the latter are smaller, refractile, and associated with a yellow to brown pigment, while areas with PAS positive diastase resistant fine cytoplasmic granules will be found in the former. Conventional cystadenocarcinoma differ from LGCCC by the lack of intraductal proliferation, golden brown pigment, solid cellular foci, and overall resemblance to atypical hyperplasia or carcinoma-in-situ of the breast. Cystadenocarcinoma tends to be an invasive tumour, whereas LGCCC is usually contained within cysts (790).

Prognosis and predictive factors
Treatment is complete surgical excision. Although the number of cases with follow-up is small, none of the cases, to date, have recurred. Greater experience and longer follow-up periods are necessary to substantiate the excellent prognosis.

Fig. 5.20 Low-grade cribriform cystadenocarcinoma (LGCCC) A The intraductal proliferations of LGCCC resemble benign breast ductal hyperplasia. The fenestrations formed are “floppy”, not rigid. B Calponin, a myoepithelial marker, highlights the largely intraductal nature of this neoplasm. LGCCC strongly expresses S100, in distinction to the papillocystic variant of acinic cell carcinoma. C Golden-brown pigment can be seen within the relatively bland tumour cells of LGCCC.
Mucinous adenocarcinoma

Definition
Mucinous adenocarcinoma is a rare malignant tumour composed of epithelial clusters within large pools of extracellular mucin. The mucin component usually occupies the bulk of the tumour mass.

ICD-O code 8480/3

Epidemiology
It usually arises in patients over 50 years of age. Males are affected more frequently than females (859, 1374, 1909, 1957, 2551).

Localization
The most frequently affected sites are the palate and the sublingual gland, followed by the submandibular gland and the upper lip. Occurrence in the parotid gland is rare (859).

Clinical features
The patients usually present with a slow-growing, painless swelling. However, local dull pain may be encountered in some cases. The tumour is firm and usually elevated.

Macroscopy
The tumour is nodular and ill defined. The cut surface is greyish-white, containing many cystic cavities with gelatinous contents.

Histopathology
The tumour is composed of round and irregular-shaped neoplastic epithelial cell nests or clusters floating in mucus-filled cystic cavities separated by connective fibrous strands. The tumour cells are cuboidal, columnar or irregular in shape, usually possess clear cytoplasm and darkly-stained, centrically placed nuclei. The tumour cells may have atypical nuclei, but mitotic figures are sparse. The tumour cells are arranged in solid clusters and tend to form secondary lumens or incomplete duct-like structures. Mucus-producing cells may arrange in a papillary pattern projecting into the mucous pools. Mucous acinus-like tumour islands may also be present. Both intracellular and extracellular mucin components show positive staining for periodic acid Schiff, Alcian blue and mucicarmine.

Immunoprofile
Immunocytochemically, the tumour cells express pan-keratin AE1/AE3 as well as cytokeratins 7, 8, 18 and 19 that are usually found in simple epithelia (859, 1374). Expression of cytokeratins 4 and 13 is seen in about 10-20%. Negative staining is noted for cytokeratins 5/6, 10, 14, 17 and for smooth muscle actin (SMA).

Electron microscopy
The cytoplasm of the tumour cells is densely packed with numerous low-electron-density mucous droplets, and seromucous droplets containing electron-dense dots are also seen. Tumour cells possessing mucous or seromucous droplets form a luminal structure, and they have irregularly arranged microvilli on the luminal side.

Differential diagnosis
Mucoepidermoid carcinoma, mucin-rich variant of salivary duct carcinoma and cystadenocarcinoma should be differentiated from mucinous adenocarcinoma. Mucoepidermoid carcinoma also shows extravasated mucin, but it consists of intermediate and epidermoid cells. Cystadenocarcinoma shows cystic spaces lined by epithelium. Extracellular mucin pools are not evident in acinic cell carcinoma.

Prognosis and predictive factors
Mucinous adenocarcinoma is insensitive to radiotherapy and has a propensity for local recurrence and regional lymph node metastases.
Oncocytic carcinoma

**Definition**
Oncocytic carcinoma is a proliferation of cytomorphologically malignant oncocites and adenocarcinomatous architectural phenotypes, including infiltrative qualities. These may arise de novo, but are usually seen in association with a pre-existing oncocytoma [1833]. Rarely, a benign appearing oncocytic tumour metastasizes following local recurrence [2498] and is designated carcinoma, despite the absence of malignant cellular morphology.

**ICD-O code**
8290/3

**Epidemiology**
Men are affected in two-thirds of cases. A wide age range from 25-91 years has been reported with a mean age of 62.5 years [71]. This neoplasm represents only 5% of oncocytic salivary gland tumours and less than 1% of all salivary gland tumours [922].

**Localization**
Nearly 80% involve the parotid gland, 8% the submandibular gland, with all others in minor salivary glands.

**Clinical features**
Typically there is a painless, nondescript mass in the parotid or submandibular gland. In cases of malignant transformation of a benign oncocytoma a rapid increase in size is noted after a period of slow growth. Facial nerve involvement may cause pain, paresis or neuropathy [922].

**Macroscopy**
They are firm, unencapsulated, tan to grey, unilocular or multilocular masses, occasionally with necrotic areas.

**Histopathology**
Sheets, islands and nests are composed of large, round to polyhedral cells with fine, granular, eosinophilic cytoplasm and central, round vesicular nuclei, often with prominent nucleoli [257]. Occasionally there are multinucleated cells. In some tumours there are duct-like structures of variable calibre. They are unencapsulated and often invade muscle, lymphatics and nerves. They are characterised cytologically by cellular atypia and pleomorphism. Histochemically, phosphotungstic acid-haematoxylin (PTAH) staining reveals fine, blue, cytoplasmic granulues. Other methods to demonstrate mitochondria such as the Novelli technique, cresylecht violet V, Klver-Barrera Luxol fast blue stains [2601] and antimitochondrial antibodies can also be used [2343].

**Immunoprofile**
Ki-67 immunostaining has been suggested in separating benign from malignant oncocytoma [1188]. In addition, alpha-1-antitrypsin staining has been helpful [476].

**Electron microscopy**
There are large numbers of mitochondria which are often abnormal in shape and size. Intracytoplasmic lumina lined with microvilli and lipid droplets have also been reported. A nearly continuous basal lamina, evenly spaced desmosomes and rearrangement of mitochondrial cristae have been demonstrated [218].

**Prognosis and predictive factors**
These high-grade tumours are characterised by multiple local recurrences and regional or distant metastases [922,940]. In one series, 7 of 11 patients studied ultimately developed metastatic disease [1227]. It appears that the most important prognostic indicator is the presence or absence of distant metastases [1833].

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**Fig. 5.23** Oncocytic carcinoma. **A** Invasion into the parotid gland. **B** Atypical tumour cells have prominent nucleoli and eosinophilic, granular cytoplasm. **C** Perineural invasion.

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Oncocytic carcinoma 235
Salivary duct carcinoma

Definition
An aggressive adenocarcinoma which resembles high-grade breast ductal carcinoma.

ICD-O code 8500/3

Synonyms
Cribriform salivary carcinoma of excretory ducts, high-grade salivary duct carcinoma

Epidemiology
Salivary duct carcinoma (SDC) is an uncommon, but not a rare form of salivary malignancy. De novo and/or expleomorphic adenoma, SDC represents 9% of salivary malignancies. The male:female ratio is at least 4:1. Most patients present after age 50 [135, 259, 1488]. The parotid is most commonly involved, but submandibular, sublingual, minor salivary gland, maxillary and laryngeal tumours have been reported [682, 745, 1383, 2021, 2583, 2862, 2909].

Etiology
A unique case of SDC arising in a longstanding chronic obstructive sialadenitis has been reported [1113].

Clinical features
Patients with SDC typically present with recent onset of a rapidly growing tumour that may fluctuate in size. Occasional patients have longer clinical histories. Pain and facial paresis may be present.

Macroscopy
SDC are usually firm, solid, tan, white or grey, with a cystic component. Infiltration of the adjacent parenchyma is usually obvious, but occasional tumours may appear to be circumscribed. SDC may also arise as the malignant component of a carcinoma ex pleomorphic adenoma, so that the macroscopic features of pleomorphic adenoma may also be present.

Tumour spread and staging
For SDC, perineural spread (60%) and intravascular tumour emboli (31%) are common. Most patients present with Stage III or IV disease, as lymph nodes are positive in 59% of patients [135].

Histopathology
SDC resembles intraductal and infiltrating mammary duct carcinoma, both architecturally and cytologically. The diagnostic “ductal lesion” comprises pleomorphic, epithelioid tumour cells with a cribriform growth pattern, “Roman bridge” formation, and intraductal comedonecrosis. The tumour infiltrates and metastasizes with a cribriform pattern, or it totally recapitulates the intrasialodochal “ductal lesion”. Solid and papillary areas may be seen, with psammoma bodies, as well as evidence of squamous differentiation. Cytologically, these cells have abundant, pink cytoplasm and large pleomorphic nuclei with prominent nucleoli and coarse chromatin. The cytoplasm may also be densely eosinophilic, granular, or oncocytic. Mitotic figures are usually abundant. Goblet cells are not seen. Rare tumours may have a prominent spindle cell or sarcomatoid growth pattern similar to the metaplastic ductal carcinomas of the breast [1064, 1819]. The mucin-rich SDC is a recently described variant of SDC [2371]. The tumour is composed of areas of typical SDC, but in addition, contains mucin lakes with islands of carcinoma cells. Another variant showing an invasive micropapillary component has also been reported [1820].

Immunoprofile
SDC is immunoreactive for low- and high-molecular-weight cytokeratin, and markers such as carcinoembryonic antigen (CEA), LeuM1, and epithelial membrane antigen (EMA) [579, 1488]. Strong nuclear reactivity for androgen receptors (AR) is reported in all SDC [1265, 1488, 2371]. SDC cells are focally positive for

Fig. 5.24 Salivary duct carcinoma. A Note the cribriform ductal component with comedonecrosis and the invasive desmoplastic carcinoma on the right. B Perineural invasion.
apocrine marker GCDFP-15 and mitochondrial antigen (MIA), and typically negative for S-100 protein, myoepithelial markers, estrogen and progesterone receptors. Variable expression of prostatic markers (prostate specific antigen, prostatic acid phosphatase) is seen (555). The MIB1 proliferative index is high, with an average value of 43% (range 25-80%). Most SDC show positive distinct membrane staining for HER-2/neu protein (1644,2392,2393).

**Differential diagnosis**

Other diagnoses to consider for SDC include metastatic breast and squamous carcinomas, oncocytic carcinoma and mucoepidermoid carcinoma. Despite a superficial resemblance to squamous carcinoma, this diagnosis can be discarded as soon as the infiltrating cribriform pattern is recognized. Identification of sialodochodysplasia supports a primary parotid origin. Goblet cells are not seen with SDC (aside from intraductal goblet cell metaplasia), thus ruling out mucoepidermoid carcinoma.

**Genetics**

Only two studies of these tumours have been published. Seven of eight tumours had LOH in at least one marker on chromosome 9p21 (351) in one study. In the other study, a high incidence of LOH was found at 6q, 16q, 17p and 17q regions (1110). Amplification of HER-2/neu gene and gene product overexpression are reported in SDC (725,1644,1803,2392,2393). Mutations and overexpression of the TP53 gene and protein are frequent (1110,1803,1823). Loss of heterozygosity at microsatellite loci, TP53 point mutations and frequent alterations of certain loci on chromosome arm 6q have been reported (1110). The chromosomal locus 9q21 contains the CDKN2A/p16 tumour suppressor gene that has been implicated in a variety of tumour types, including SDC (351). More polymorphic genetic markers located at this particular region suggest that inactivation of CDKN2A/p16 gene is associated with progression of SDC (351).

**Prognosis and predictive factors**

SDC is one of the most aggressive salivary malignancies. A review of 104 cases concluded that 33% of patients developed local recurrence and 46% developed distant metastasis (135). Sites for distant metastasis include lungs, bones, liver, brain and skin. Sixty-five percent of patients died of disease, between 5 months to 10 years, usually within 4 years of diagnosis. The clinical course is characterized by early distant metastases. Tumour size, distant metastasis, and HER-2/neu overexpression are putative prognostic parameters for SDC, while expression of p53 protein, DNA aneuploidy, and proliferative activity do not correlate with outcome (2393). The clinical outcome for the mucin-rich variant of SDC is similar to that of conventional SDC (2371). The invasive micropapillary variant appears to be particularly aggressive (1820).
Adenocarcinoma, not otherwise specified

Definition
Adenocarcinoma, not otherwise specified, is a malignant salivary gland tumour that exhibits ductal differentiation but lacks any of the histomorphologic features that characterize the other defined types of salivary carcinoma. The modifying term “not otherwise specified” should be included because most other epithelial salivary gland malignancies are also adenocarcinomas.

ICD-O code 8140/3

Synonyms
These tumours have often been reported as miscellaneous or unclassified adenocarcinomas or, simply, as adenocarcinoma [1662,1815,2447]. It appears that many reports include cases that should be classified as one of the more specific carcinoma types [668]. They should not be grouped together with tumours that arise from the seromucous glands of the nasal cavity, paranasal sinuses or larynx because in these sites they appear to have a more aggressive biologic behaviour [2447].

Epidemiology
The inconsistent reporting of these tumours limits our understanding of them. In one report they are second in frequency only to mucoepidermoid carcinoma among malignant salivary gland tumours and account for about 17% of the carcinomas [668]. Women outnumber men slightly and the average patient age is 58 years. They are extremely rare in children.

Localization
About 60% and 40%, respectively, occur in the major and minor glands. The vast majority that involve the major glands occur in the parotid, and the minor gland tumours most often arise from the glands in the hard palate, buccal mucosa, and lips.

Clinical features
Most patients with tumours of major glands present with solitary, asymptomatic masses, but about 20% have pain or facial weakness [2447]. Pain is more often associated with tumours of the submandibular glands. Minor gland tumours may be ulcerated and about 25% of palatal tumours involve the underlying bone. Tumour duration ranges from one to 10 years [2447].

Macroscopy
Adenocarcinoma, NOS, is often partially circumscribed but in many areas the periphery is irregular and ill defined. Areas of necrosis or haemorrhage may contrast with the white or yellowish cut surface.

Histopathology
Shared by all tumours in this group are the presence of glandular or duct-like structures, infiltrative growth into parenchyma or surrounding tissues, and lack of features that characterize other salivary adenocarcinomas. There is considerable variability in the architectural structure. Some have small confluent nests or cords of tumour cells, others large discrete islands with intervening trabeculae of fibrous connective tissue, and still others large solid, densely cellular sheets. This latter group reveals very limited stromal connective tissue.

Ductal differentiation is widespread in low and intermediate grade tumours but usually much more subtle in high-grade tumours. Small cysts are occasionally present in those with numerous ducts. Cuboidal or ovoid cells predominate in most tumours but scattered clear and oncocytic cells are occasionally evident. Small deposits of eosinophilic acellular material and extracellular mucin may be present. Unlike most other salivary adenocarcinomas, the cytologic variability is useful for grading these tumours [2447]. Low-grade

Fig. 5.27 Adenocarcinoma, not otherwise specified. A Architectural and cellular variability. Prominent ductal differentiation is present, within closely arranged tumour islands. B Focal tubular structures with hyaline cores, reminiscent of adenoid cystic carcinoma.
Adenocarcinoma, not otherwise specified

Tumours demonstrate minimal variability of nuclear size, shape, or staining density, and rare mitoses. In some, the bland nuclear morphology suggests benignity and determination of their malignant nature is based largely on the identification of invasive growth. Intermediate grade tumours show nuclear variability and more frequent mitoses. High-grade tumours have enlarged, pleomorphic, hyperchromatic nuclei, focal necrosis, and frequent and atypical mitoses. The presence of ductal differentiation helps in the distinction from undifferentiated carcinoma.

**Differential diagnosis**

Because these tumours do not have pathognomonic histopathologic features, the possibility of metastatic adenocarcinoma should be considered. While immunohistochemical studies may be useful in this evaluation [2370] it should be remembered that immunoreactivity with prostate-specific antigen has been reported [2571, 2574].

**Prognosis and predictive factors**

Limited data suggest that the clinical stage, site of involvement and grade of tumour influence prognosis [1662, 2447, 2708]. Minor gland tumours have a better prognosis than those of the major glands. Distant metastases may occur despite regional control and recurrence is more frequent with high-grade tumours [2447]. In one study, the 15-year survival for low, intermediate and high-grade tumours was 54, 31, and 3%, respectively, and the cure rate of the low-grade tumours was similar to that of acinic cell adenocarcinoma [2447].
Definition
Myoepithelial carcinoma of the salivary glands is a neoplasm composed almost exclusively of tumour cells with myoepithelial differentiation, characterized by infiltrative growth and potential for metastasis. This tumour represents the malignant counterpart of benign myoepithelioma.

ICD-O code 8982/3

Synonym
Malignant myoepithelioma

Epidemiology
The mean age of patients at presentation is 55 years with a wide age distribution (range 14-86). Males and females are affected equally. In large series, myoepithelial carcinomas comprise less than two percent of all salivary gland carcinomas, but they may not be as rare as has been suggested before [2251,2304]. The very low historic incidence is probably due to their recent recognition as a separate tumour entity.

Etiology
No etiological factors are known.

Localization
Most cases (75%) arise in the parotid, but they also occur in the submandibular and minor glands.

Clinical features
The tumours are locally destructive. The majority of patients present with the complaint of a painless mass.

Macroscopy
Myoepithelial carcinomas are unencapsulated but may be well-defined with nodular surfaces. Tumour size varies considerably (2-10 cm). The cut surface is grey-white and can be glassy. Some tumours show areas of necrosis and cystic degeneration.

Tumour spread and staging
They can involve adjacent bone. Perineural and vascular invasion may occur. Regional and distant metastases are uncommon at presentation, but may occur late in the course of disease.

Histopathology
Myoepithelial carcinoma characteristically has a multilobulated architecture. The range of cell types in myoepithelial carcinoma reflects that seen in its benign counterpart. The tumour cells often are spindled, stellate, epithelioid, plasmacytoid (hyaline), or, occasionally, vacuolated with signet ring like appearance. Other tumours tend to be more cellular composed of spindle-shaped cells, and they can resemble sarcoma. Rarely, myoepithelial carcinoma is composed of a monomorphic population of clear cells with myoepithelial features [1719]. The tumour cells may form solid and sheet-like formations, trabecular or reticular patterns, but they can also be dissociated, often within plentiful myxoid or hyaline stroma. The neoplastic nodules frequently have necrotic centres. Pseudocystic or true cystic degeneration can occur. Sparse areas with squamous differentiation may be found. Rarely, myoepithelial carcinoma contains duct-like lumina usually with non-luminal cell differentiation of the lining cells. A tumour containing more than the occasional true luminal cell should not be included in the category of purely myoepithelial neoplasia. Different cell types and architectural patterns may be found within the same tumour. In fact, most myoepithelial carcinomas are less monomorphic than benign myoepithelioma. They also may demonstrate high mitotic activity with considerable variation [595,1154,1827,2251]. Cellular pleomorphism can be marked, and necrosis may occur [1827,2251]. However, unequivocal evidence of infiltrative, destructive growth is the major requirement for diagnosis, and it is this property that distinguishes myoepithelial carcinoma from benign myoepithelial tumours.

Immunoprofile
Reactivity for cytokeratin and at least one other myoepithelial marker is always present. A monoclonal antibody for myoepithelial differentiation is also useful for this purpose.

Fig. 5.29 Myoepithelial carcinoma. A Multinodular growth pattern. B Spindle cell malignant myoepithelioma invading soft tissues.
of the other myoepithelial markers, including smooth muscle actin, GFAP, CD10, calponin and smooth muscle myosin heavy chain, is required for diagnosis [595,1827].

**Electron microscopy**
Ultrastructural criteria for the diagnosis of myoepithelial carcinoma include longitudinally oriented 6-8 nm fine cytoplasmic microfilaments with focal dense bodies, pinocytic vesicles, desmosomes and hemidesmosomes, basal lamina and intermediate filaments [41,640].

**Precursor lesions**
Myoepithelial carcinomas may arise de novo, but it is important to note that about half of cases develop in pre-existing pleomorphic adenomas, or from benign myoepitheliomas, particularly in recurrences [595,1827,2251].

**Genetics**
Comparative genomic hybridization has revealed infrequent abnormalities in these lesions with only three of 12 myoepitheliomas manifesting various chromosomal losses. Of myoepithelial carcinomas, five have manifested chromosome 8 alterations [1154].

**Prognosis and predictive factors**
Myoepithelial carcinomas are locally aggressive salivary gland neoplasms that exhibit diverse clinical outcomes. Approximately one third of patients die of disease, another third have recurrences, mostly multiple, and the remaining third are disease free. Marked cellular pleomorphism and high proliferative activity correlate with a poor clinical outcome [1827,2251]. There is no difference in clinical behaviour of "de novo" myoepithelial carcinomas and of those arising in pleomorphic adenomas and benign myoepitheliomas [595,2251].
Carcinoma ex pleomorphic adenoma

Definition
Carcinoma ex pleomorphic adenoma is defined as a pleomorphic adenoma from which an epithelial malignancy is derived.

ICD-O code 8941/3

Synonyms
Carcinoma arising in a benign mixed tumour, carcinoma ex benign mixed tumour, carcinoma arising in a pleomorphic adenoma, malignant mixed tumour.

Epidemiology
Many large series of carcinoma ex pleomorphic adenoma (Ca-ex-PA) have been reported and recently summarized: they comprise approximately 3.6% of all salivary tumours (range 0.9-14%), 12% of all salivary malignancies (range 2.8-42.4%), and 6.2% of all pleomorphic adenomas (range 1.9-23.3%) [898]. Ca-ex-PA usually presents in the 6th or 7th decades, approximately one decade later than patients with pleomorphic adenoma.

Etiology
Many Ca-ex-PA probably result from the accumulation of genetic instabilities in long-standing pleomorphic adenomas.

Localization
Ca-ex-PA most frequently arises in the parotid gland; but may also originate from the submandibular gland and minor salivary sites, most commonly the palate, occasionally with involvement of the nasopharynx [838].

Clinical features
The typical history is that of a long-standing mass present much longer than 3 years with rapid growth over the previous few months; however, a significant proportion of patients present with a clinical history of less than three years [898,1533]. Patients frequently complain of a painless mass; but pain, facial nerve palsy, and skin fixation may also occur.

Macroscopy
The average size of Ca-ex-PA is more than twice that of its benign counterpart, ranging from 1.5-25 cm in greatest diameter [786,2624]. Grossly, Ca-ex-PAs are usually poorly circumscribed and many are extensively infiltrative. Occasionally, tumours are well circumscribed, scar-like or appear completely encapsulated [252,2624].

Histopathology
The proportion of benign versus malignant components can be quite variable. Occasionally, extensive sampling is necessary to find the benign component and in rare cases, a benign remnant might not be found. But if there is clinicopathologic documentation of a previously excised pleomorphic adenoma in the same site, then the malignancy can also be classified as a Ca-ex-PA.

The malignant component is most frequently a poorly differentiated adenocarcinoma (salivary duct type or not otherwise specified) or an undifferentiated carcinoma; however, virtually any form of carcinoma may be found [898,1338, 1491]. An infiltrative, destructive growth pattern is the most reliable diagnostic criterion. Nuclear hyperchromasia and pleomorphism are frequent, although occasional tumours may demonstrate minimal atypia. This latter feature (tumour grade) directly correlates with prognosis. Necrosis is often present and mitoses are usually easy to find.

Ca-ex-PAs should be subclassified into non invasive, minimally invasive (≤1.5 mm penetration of the malignant component into extra capsular tissue) and invasive (>1.5 mm of invasion from the tumour capsule into adjacent tissues), as the first two groups usually have an excellent prognosis while the latter has a more guarded prognosis. The distinction between noninvasive and invasive tumours is based on destructive invasion through the capsule into peritumoral tissues. Non-invasive Ca-ex-PAs are also referred to as carcinoma in-situ arising in a pleomorphic adenoma, intracapsular carcinoma ex pleomorphic adenoma or pleomorphic adenoma with severe dysplastic changes. Atypical changes within these tumours range from focal to diffuse often

Fig. 5.32 Carcinoma ex pleomorphic adenoma. A Invasive type. Pleomorphic adenoma and carcinoma components are seen in the left and right, respectively. Carcinoma component is reminiscent of salivary duct carcinoma in this case. B Non-invasive type. Carcinoma cells replacing the inner ductal layer leaving outer benign myoepithelial layer.
with multifocal areas containing carcinoma, which frequently overgrows and replaces many of the benign elements. The earliest changes typically consist of tumour cells replacing the normal inner duct epithelial layer leaving the normal peripherally located myoepithelial layer intact.

**Differential diagnosis**
The most important differential diagnosis is between minimally invasive Ca-ex-PA and the more typical invasive Ca-ex-PA. This differential has prognostic significance, and affects decisions regarding the need for lymph node dissection and adjuvant radiotherapy. Also carcinomas may rarely arise in a histologically benign "adenoma" ("monomorphic" adenoma); they appear to have a more favourable prognosis [1576].

**Genetics**

**Cytogenetics**
Deletions of chromosome 5(q22-23, q32-33) and t(10;12) (p15;q14-15) with 12q breakpoint at the 5' of the HMGIC and translocation of the entire gene to the 10 marker chromosome followed by deletion/amplification of the segment containing HMGIC and MDM2 genes have been reported [653,1220,2193]. Rearrangements of 8q12 are a frequent finding. Alterations at 12q13-15 with amplification of HMGIC and MDM2 genes have also been reported [2125]. Cytogenetic evidence of amplification (homogeneously stained region and double minute) was found in 40% of these tumours. Both genes may contribute to the malignant transformation of pleomorphic adenoma. Alterations at chromosomes 6q deletion and 8q rearrangements have been reported.

**Molecular genetics**
Microsatellite analysis of these tumours has shown LOH at chromosome 8q and 17p. Concurrent analysis of the benign and malignant components of these tumours showed 8q and/or 12q in both components and additional alterations in 17p only in the carcinoma [651]. In another study homozygous deletion of the p16 gene on chromosome 9p21 was found in carcinoma of one case and microsatellite instability was noted in both the adenoma and carcinoma components in two tumours [2510]. A single case report of a carcinosarcoma, in which the carcinoma and the sarcoma components were concurrently analyzed, showed lack of p53 alterations and concomitant LOH at different loci on chromosome 17 and 18 supporting monoclonality [932].

**Prognosis and predictive factors**
In general, the recommended therapy is wide local excision with contiguous lymph node dissection. Adjuvant radiation therapy is recommended for widely invasive tumours. If the carcinomatous component is low-grade and/or minimally invasive and if the tumour is adequately excised, then adjuvant radiation therapy may not be necessary. Patients with non-invasive or minimally invasive Ca-ex-PA typically have an excellent prognosis, similar to benign pleomorphic adenoma. Metastatic spread is exceptional [726]. Invasive Ca-ex-PAs, as a group, are extremely aggressive malignancies with approximately 23-50% of patients developing one or more recurrences [786, 898,1491,1533]. The metastatic rate varies with each series; up to 70% of patients develop local or distant metastasis [877,898,1491]. The metastatic sites in order of frequency are lung, bone (especially spine), abdomen and central nervous system [786,2592]. Ca-ex-PA with capsular penetration of more than 1.5 mm is associated with a poor prognosis; survival rates at 5, 10, 15, and 20 years range from 25-65%, 18-50%, 10-35%, and 0-38%, respectively [786,877,1491,1533,2592,2624]. Therefore, it is important to designate those Ca-ex-PA that are confined within the capsule and those invading through the capsule as non-invasive or invasive, respectively, and to differentiate within the latter group between widely invasive and minimally invasive tumours.

One study showed that no patient with less than 8 mm invasion from the capsule died from the tumour, whereas all patients with invasion greater than 8 mm beyond the capsule ultimately died of disease [2624]. The local recurrence rate (LRR) in this latter series also correlated with extent of invasion; a LRR of 70.5% was found for tumours with invasion beyond 6 mm from the capsule, as compared to a LRR of 16.6% for tumours with invasion of less than 6 mm. In another study consisting of four patients with 5 mm of invasion beyond the tumour capsule, two died of disease and two were alive and well [1491]. The two patients with less than 5 mm of invasion (2 and 3 mm) were alive and well with no evidence of disease. Also, all four patients with intracapsular carcinoma were alive and well without evidence of disease progression.

The improved prognosis for minimally invasive tumours has been confirmed by Brandwein et al who observed recurrence free for periods ranging from 1-4 years (mean 2.5 years) [252]. Tumour size and grade are also significant prognosticators in the more widely invasive Ca-ex-PAs. The five-year survival rates have been correlated with histologic subtype of the carcinoma component: there was a 30% survival rate for undifferentiated carcinomas, 50% for myoepithelial carcinomas, 62% for ductal carcinomas and 96% survival rate for terminal duct carcinomas [2624]. In addition, 63% of patients with high-grade carcinomatous components died of the disease, while patients with lower grade carcinomatous elements did not [1491].
Carcinosarcoma

Definition
Carcinosarcoma is a malignant tumour composed of a mixture of both carcinomaous and sarcomatous elements.

ICD-O code 8980/3

Synonym
True malignant mixed tumour

Epidemiology
Carcinosarcoma is extremely rare; approximately 50-60 cases have been reported to date [47,898,911,932,1010,1320,2377,2466]. The mean age at presentation was 58 years with a range of 14-87 years [899]. A number of patients have had a history of recurrent pleomorphic adenoma [2466] and several cases have arisen in a pleomorphic adenoma (carcinosarcoma ex pleomorphic adenoma) [899,1010,1320,1400].

Localization
Two-thirds have arisen in the parotid gland, approximately 19% in the submandibular glands, and 14% in the palate [899]. One case has been reported in the tongue and one in the supraglottic region [691].

Clinical features
Patients typically present with a mass, which may be painful.

Macroscopy
Tumours are well to poorly circumscribed.

Histopathology
The tumour is composed of mixtures of carcinomaous and sarcomatous elements in varying proportions [225]. Chondrosarcoma and osteosarcoma are the most common sarcomatous elements and moderate to poorly differentiated ductal carcinoma or undifferentiated carcinoma are the most common carcinomatous components. Local tissue infiltration and destruction are characteristic of this neoplasm.

Genetics
LOH at 17p13.1, 17q21.3 and 18q21.3 has been found in one carcinosarcoma. Sequencing studies excluded TP53 mutations, suggesting inactivation of another tumour suppressor gene at 17p13 [932].

Prognosis and predictive factors
Treatment is wide surgical excision combined with radiotherapy. Almost 60% of patients die of local recurrence and/or metastatic disease (lungs, bones, central nervous system), usually within a thirty month period [47,899,2466].

Fig. 5.34 Carcinosarcoma. Low power. The majority of this tumour is composed of poorly differentiated sarcoma with focal areas of poorly differentiated adenocarcinoma at the periphery.

Fig. 5.35 Carcinosarcoma A Tumour composed of mixtures of adenocarcinomatous and osteosarcomatous components. B Midportion of tumour. Note areas with chondrosarcomatous differentiation (right side) and a small focus with osteosarcomatous differentiation (upper left).
Metastasizing pleomorphic adenoma

**Definition**
A histologically benign pleomorphic adenoma that inexplicably manifests local or distant metastasis.

**ICD-O code**
8940/1

**Synonyms**
Metastasizing benign mixed tumour, malignant mixed tumour.

**Epidemiology**
To date, approximately 40 cases have been described (406,2768).

**Etiology**
It has been postulated that multiple recurrences and surgical procedures allow some tumours to gain venous access and metastasize.

**Localization**
Greater than three-quarters arise in the parotid gland, 13% in the submandibular gland and 9% in the palate.

**Histopathology**
Characteristically, the primary salivary gland tumour and metastases are composed of the typical mixture of benign-appearing epithelial and mesenchymal components of a pleomorphic adenoma. The histology is not predictive regarding its ability to metastasize. Mitotic figures and nuclear pleomorphism may be seen, but the tumour is not overtly histologically malignant.

**Prognosis and predictive factors**
The treatment of choice is surgical excision. Metastasizing pleomorphic adenomas are characterized by multiple local recurrences and a long interval (1.5-55 years) between development of the primary tumour and its metastasis. Half of the tumours metastasize to bone, 30% to lung and 30% to lymph nodes; rarely tumours spread to other body sites. Forty percent of patients died with disease; 47% were alive and well, and 13% were alive with disease (899).

Squamous cell carcinoma

**Definition**
A primary malignant epithelial tumour composed of epidermoid cells, which produce keratin and/or demonstrate intercellular bridges by light microscopy. It is essential to exclude the possibility of metastatic disease. By convention, the diagnosis of salivary squamous cell carcinoma is restricted to the major salivary glands, since minor salivary squamous carcinomas cannot be reliably distinguished from tumours of mucosal origin.

**ICD-O code**
8070/3

**Synonym**
Epidermoid carcinoma

**Epidemiology**
Primary squamous cell carcinoma (PSCC) probably represents less than 1% of salivary gland tumours. PSCC occurs in patients over a wide age range, but the majority present in the 6th through 8th decades, with a mean of 60-65 years. They are unusual in patients younger than 20 years, although several cases have been described in children (669). There is a male to female ratio of approximately 2:1.

**Etiology**
In several studies, PSCC has been associated with a history of prior radiotherapy, with a latent period of 15-30 years (2329).

**Localization**
Roughly 80% of PSCC arise in the parotid gland and 20% in the submandibular gland. PSCC of the sublingual gland is quite unusual. Occasionally, cases arise from the mucosa lining Stensen's duct.

**Clinical features**
Patients with PSCC present with a rapidly enlarging mass, which is frequently painful. Tumours are firm and fixed and may be associated with facial nerve weakness. PSCC is typically high stage at the time of diagnosis (2329,2468).

**Macroscopy**
PSCC is an invasive neoplasm with ill-defined margins. Most tumours are greater than 3 cm in size. The cut surface is typically solid, firm, and light grey or tan to white, sometimes with focal necrosis.
Histopathology

The histology of PSCC of salivary origin is similar to that of well- to moderately-differentiated squamous cell carcinoma originating elsewhere in the head and neck. The tumour infiltrates the salivary parenchyma in irregular nests and trabeculae, accompanied by a fibrous to desmoplastic stromal response. Squamous metaplasia and dysplasia of salivary ducts are occasionally identified in association with PSCC. Perineural invasion and extension into adjacent soft tissue are common findings. There is a significant incidence of cervical nodal metastases (both clinically apparent and occult) at the time of initial surgery (779, 869, 1456, 2329).

Differential diagnosis

The most critical distinction in the differential diagnosis of PSCC is ruling out the possibility of metastatic squamous cell carcinoma, whose incidence is greater than true PSCC. PSCC must also be distinguished from mucoepidermoid carcinoma (MEC). MEC is typically composed of a variable cell population, including mucocytes, basaloid, and intermediate cells, in addition to epidermoid cells. However, prominent keratinization is not characteristic of MEC. MEC may exhibit cystic areas and focal clear cell differentiation, features not observed in PSCC. Histochemical stains for intracellular mucin to rule out high-grade MEC are recommended before making a definitive diagnosis of PSCC (669). Squamous metaplasia in infarcted or surgically manipulated tumours can be misinterpreted as PSCC.

Keratocystoma is a recently described, rare lesion of salivary glands that may be confused with squamous cell carcinoma (1822). It is characterized by multilocular spaces lined by stratified squamous cells containing keratotic lamellae and focal solid epithelial nests. The consistent absence of metastasis, necrosis or invasion, as well as the lack of cytological atypia and minimal cellular proliferative activity in keratocystoma is essential in distinguishing this lesion from PSCC.

Genetics

Cytogenetic studies in several cases of PSCC have yielded somewhat variable results, although it appears that various 6q deletions may be common, similar to the findings in other salivary carcinomas (1222). Interestingly, this karyotype is unusual in squamous cell carcinoma of other head and neck sites (1222).

Prognosis and predictive factors

PSCC is considered a relatively high-grade, aggressive salivary carcinoma. Five-year disease specific survival is approximately 25-30%. Local-regional recurrence develops in at least half of patients and distant metastases are found in 20-30% (2329). Overall, 75% die of their disease, usually within 5 years (1456, 2329). In the largest published specific analysis of PSCC (2329), tumour stage was the most important prognostic factor. Age greater than 60 years, ulceration, and fixation also had a significant negative impact on survival. Two additional series, which only considered parotid tumours, reported that age, facial nerve paralysis, deep fixation, and type of treatment were of statistical significance (869, 1456).
**Small cell carcinoma**

**Definition**
Small cell carcinomas of the salivary glands are rare, malignant epithelial tumours characterized by a proliferation of small anaplastic cells with scant cytoplasm, fine nuclear chromatin, and inconspicuous nucleoli.

**ICD-O code** 8041/3

**Synonyms**
Small cell undifferentiated carcinoma, small cell anaplastic carcinoma, oat cell carcinoma, neuroendocrine carcinoma.

**Epidemiology**
They account for less than 1% of all salivary gland tumours and approximately 2% of salivary gland malignancies [668]. Most patients are older than 50 years at the time of initial diagnosis; however, these tumours have been described in younger patients [668,902]. The tumour has a slight predilection for males.

**Localization**
The tumours can involve major and intraoral minor salivary glands, and are most common in the parotid gland.

**Clinical features**
Patients typically present with a painless, rapidly growing mass of several months duration. Cervical lymphadenopathy and facial nerve palsy are common findings. Paraneoplastic syndromes accompanied by the production of ectopic hormones are unusual [1746].

**Macroscopy**
It is a firm, poorly circumscribed tumour that often infiltrates the surrounding salivary gland parenchyma and adjacent soft tissues. The tumour is usually grey to white and commonly accompanied by necrosis and haemorrhage.

**Histopathology**
Small cell carcinoma is characterized by sheets, cords, or irregular nests of anaplastic cells and a variable amount of fibrous stroma. The tumour cell nests may exhibit a peripheral palisading pattern. Rosette-like structures are occasionally seen. Tumour cells are usually 2-3 times larger than mature small lymphocytes and have round to oval nuclei with scant cytoplasm. Fusiform or polygonal cells as well as occasional larger cells are sometimes observed. Nuclear chromatin is finely granular, and nucleoli are absent or inconspicuous. Cell borders are ill defined, and nuclear moulding is common. Mitotic figures are numerous. A tumour may have small foci of ductal differentiation [902]. Focal areas of squamous differentiation also have been described [1030,2196]. Extensive necrosis and vascular and perineural invasion are common.

**Immunoprofile**
In most small cell carcinomas, the tumour cells express at least one neuroendocrine marker such as chromogranin A, synaptophysin, CD57 (Leu-7), CD56 (neural cell adhesion molecule) and neurofilament [907,1818]. However,
immunoreactivity for neuron-specific enolase alone is insufficient evidence for confirming the neuroendocrine differentiation of the tumour. Most small cell carcinomas are positive for cytokeratins, which often have a characteristic paranuclear dotlike pattern of reactivity \cite{372,1818}. The majority of the tumours are also positive for epithelial membrane antigen \cite{907,1818}. Similar to Merkel cell carcinoma, but unlike pulmonary small cell carcinoma, three out of four salivary small cell carcinomas are cytokeratin 20 positive \cite{1818}. Also, small cell carcinomas are negative for S-100 protein and HMB-45.

**Electron microscopy**

Electron microscopic examination shows membrane-bound neuroendocrine granules in about one-third of small cell carcinomas \cite{907}. The tumour cells contain sparse cytoplasmic organelles, and either poorly or well-formed desmosomes interconnect the cells. Multidirectional differentiation with the presence of myofilament-like microfilaments and tonofilaments has been reported \cite{1030,2628,2836}.

**Prognosis and predictive factors**

Local recurrence and distant metastases develop in more than 50% of patients after the initial diagnosis. Cervical lymph node involvement is less common than haematogenous metastasis. The 5-year survival rate for patients with small cell carcinomas arising in the major salivary glands ranges from 13 to 46\% \cite{902,1818,2042}. Overall survival is reduced for patients with a primary tumour larger than 3 cm, negative immunostaining for cytokeratin 20 and decreased immunoreactivity for neuroendocrine markers \cite{1818}.

![Fig. 5.41 Small cell carcinoma. High-power view showing tumour cells with scant cytoplasm, finely granular nuclear chromatin, and inconspicuous nucleoli. Mitotic figures are readily identified (A, B). A Tumour cell nuclei are oval to spindle, with dense chromatin. B The tumour cells are slightly larger than A and they have rather pale, dispersed chromatin and a little more abundant cytoplasm. C Tumour cells are diffusely immunopositive for chromogranin A. D Paranuclear dotlike pattern of immunoreactivity for cytokeratin 20.](image)
Large cell carcinoma

Definition
Large cell carcinomas are rare, high-grade malignant salivary gland epithelial tumours composed of pleomorphic cells with abundant cytoplasm and absence of features of other specific tumour types.

ICD-O code 8012/3

Synonym
Large cell undifferentiated carcinoma.

Epidemiology
Large cell carcinomas are exceptionally rare [1151,1816]. In the majority of cases, the patients were older than 60 years. Males and females are affected equally.

Localization
The majority of large cell carcinomas arise in the major salivary glands, especially the parotid gland [1151,1432,1768,1816,1828,2836]. A few tumours of minor salivary gland origin have been reported [1768].

Clinical features
Many patients present with a rapidly growing firm mass that often is fixed to adjacent tissue. Facial nerve paralysis and cervical lymph node enlargement are common findings.

Macroscopy
A large cell carcinoma is usually a poorly circumscribed, solid tumour with greyish white or tan cut surface. Necrosis and haemorrhage are easily found. Invasion into the adipose and muscular tissue adjacent to the salivary gland is common.

Histopathology
The tumour is composed of large, pleomorphic cells (>30µm) with an abundance of eosinophilic or occasionally clear, cytoplasm. In some tumours there is striking dyscohesive architecture resembling lymphoma. The tumour cell nuclei have a polygonal or fusiform shape, prominent nucleoli, and coarse chromatin with a vesicular distribution. Cell borders are usually well-defined. Bizarre giant tumour cells may be present. Mitotic figures are readily identified. The tumour growth pattern consists of sheets and trabeculae, with a conspicuous tendency for necrosis. Organoid, rosette-like, and peripheral palisading patterns characterize some of the large cell carcinomas [1828]. Rare foci of ductal or squamous differentiation can be present in large cell carcinomas. Lymphoid cell infiltration is usually focal and patchy. Perineural and vascular involvement is prominent.

Immunoprofile
Some cases of large cell carcinoma may be positive for one of the neuroendocrine markers, including chromogranin A, synaptophysin, CD57 (Leu-7), PGP9.5, or CD56 (neural cell-adhesion molecule). No immunoreactivity for cytokeratin 20 was found. The Ki-67 (MIB-1) labeling index is high and often greater than 50%. In two reported cases, the tumour cells showed diffuse immunoreexpression of bcl-2 protein, epidermal growth factor receptor, and cyclin D1 and reduced immunoexpression of p21/waf1 and p27/kip1 [1828]. Diffuse TP53 nuclear immunoexpression has been found in 4 of 5 cases [1803,1828, 2421].

Fig. 5.42 Large cell carcinoma. A Sheet-like growth pattern of large pleomorphic cells with abundant eosinophilic cytoplasm and prominent nucleoli. B Strong immunoreactivity for cytokeratin.
Electron microscopy
Ultrastructurally, tumour cells occasionally have a squamous or glandular differentiation not apparent on conventional light microscopic examination (1816, 2836). Neuroendocrine differentiation is rare; neurosecretory granules have been described in 3 cases (1151, 1432, 1828). Prominent desmosome-like junctions connect the tumour cells.

Genetics
Genetic studies of salivary gland large cell carcinoma are scant. TP53 mutation has been detected in two of three cases, and 1 case demonstrated loss of heterozygosity (LOH) at chromosome 17p (1803, 1828). Two cases of large cell neuroendocrine carcinoma exhibited LOH at chromosome 9p21 (1828).

Prognosis and predictive factors
Large cell carcinoma is an aggressive tumour with a propensity for local recurrence, cervical lymph node metastases, and distant spread. However, one study has shown that cell size (small vs large type of carcinoma) has no influence on prognosis (1151). Tumour size has been found to be a prognostic indicator; all patients with tumours larger than 4 cm died of disease with distant metastases (1151).

Fig. 5.43 Large cell carcinoma. A Organoid growth pattern. B Solid growth with peripheral palisading and rosette-like structures. Tumour cells have large and polygonal nuclei with vesicular chromatin and prominent nucleoli.
Lymphoepithelial carcinoma

**Definition**
Lymphoepithelial carcinoma (LEC) is an undifferentiated carcinoma accompanied by a prominent non-neoplastic lymphoplasmacytic infiltrate.

**ICD-O code** 8082/3

**Synonyms**
Lymphoepithelioma-like carcinoma (LEC) {1173,1387}; malignant lymphoepithelial lesion (236,2253); undifferentiated carcinoma with lymphoid stroma (459,2304); undifferentiated carcinoma (986,1359); carcinoma ex lymphoepithelial lesion (152).

**Epidemiology**
LEC of the salivary gland is rare, accounting for less than 1% of all salivary gland tumours. It shows a striking racial predilection for Inuits (Eskimo) in the Arctic regions (Greenland, Canada, Alaska), South-eastern Chinese, and Japanese {32,236,986,1479,1821,2253,2326}. The Inuit populations have the highest worldwide incidence of malignant salivary gland tumours, with the majority represented by LEC {32,236,1708}. Slight female predominance, higher frequency of parotid gland involvement, more frequent high stage disease and apparently more aggressive clinical course have been reported in Inuits [236,1428,1479,1708,2253,2326,2636]. Patients affected by LEC span a wide age range from the first to the ninth decades, with most cases occurring in the fifth decade. There is a slight male predominance {236}.

**Etiology**
The near 100% association of Epstein-Barr virus (EBV) with salivary gland LEC from the endemic areas, and the presence of the virus in a clonal episcopal form suggest an important role of EBV in tumourigenesis {31,236,986,1143,1387,1428,1479,1821,2636}. Serologic studies show elevated titres of anti-EBV viral capsid antigen IgA or anti-EBV nuclear antigen IgG, though non-specific, in more than 50% of patients with salivary gland LEC from the endemic areas {31,236,986,1143,1387,1428,1479,1821,2636}. In patients from non-endemic areas, EBV is usually absent, although rare cases may harbour the virus {209,857,1173,1359}. These findings indicate complex interactions of ethnic, geographic and viral factors in the pathogenesis of salivary gland LEC.

**Localization**
The parotid gland is affected in approximately 80% of the cases, followed by the submandibular gland {1479,2253,2326,2636}. LEC can also rarely occur in the minor salivary glands of the oral cavity, oropharynx and hypopharynx.

**Clinical features**
LEC presents as a parotid or submandibular swelling (which may be long-standing with recent rapid increase in size), with or without pain {236,857,2253}. Advanced tumours may become fixed to the underlying tissues or the skin, although facial nerve palsy occurs in only about 20% of cases. Cervical lymph node involvement, which may be extensive, is seen in 10-40% of cases at presentation {236,1000,1387,1479,2253,2636}. There is no clinical or serologic evidence of an underlying Sjögren syndrome {1373,1479,2253}. Since LEC of salivary gland is morphologically indistinguishable from nasopharyngeal carcinoma (which is much more common), it is important to examine and biopsy the nasopharynx thoroughly before accepting the salivary gland tumour as primary LEC {377,2252}.

**Macroscopy**
The tumours can be circumscribed or show frank invasion into the surrounding tissues which include lymphoid follicles (blue-staining).

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**Fig. 5.44** Lymphoepithelial carcinoma of parotid gland. In this example, irregular islands of carcinoma (purple-staining) are intermingled with abundant lymphoid tissues which include lymphoid follicles (blue-staining).
gland and extraglandular soft tissues. They are fleshy and firm, and range from 1-10 cm in size (mean 2-3 cm) [2252].

Tumour spread and staging
LEC has a propensity to spread to regional cervical lymph nodes [236, 1479,2636]. Distant metastasis, which can be found in up to 20% of cases at presentation, tends to occur in the lung, liver, bone and brain. In metastatic deposits, the prominent lymphoplasmacytic infiltrate characteristic of the primary lesion may or may not be present.

Histopathology
The tumour grows in infiltrative sheets, islands and cords separated by a lymphoïd stroma. The tumour cells possess indistinct cell borders, lightly eosinophilic cytoplasm, oval vesicular nuclei with open chromatin, and conspicuous nucleoli. The nuclei usually show moderate variation in size, although rare cases exhibit fairly uniform-appearing nuclei. Necrosis and mitotic figures are usually easily found. Sometimes the tumour cells can be plump and spindly, with formation of fascicles [445]. Focal squamous differentiation in the form of increased amount of eosinophilic cytoplasm and vague intracellular bridges is occasionally present.

The tumour is by definition richly infiltrated by lymphocytes and plasma cells, often accompanied by reactive lymphoid follicles. The lymphoid component can sometimes be so heavy that the epithelial nature of the tumour may not be evident. Histioocytes are abundant in the tumour islands in some cases, imparting a “starry sky” appearance [2253]. Other inconsistent findings are non-caseating granulomas with or without multinucleated giant cells, amyloid deposition [1387], cyst formation in some tumour islands, perineural and lymphovascular invasion. Tumour cells are immunoreactive for pan-cytokeratin and epithelial membrane antigen. The lymphoid cells include a mixture of B cells and T cells. Electron microscopy shows features of squamous differentiation, with desmosomes and tonofilaments.

In endemic cases, EBV-encoded RNA (EBER) and EBV-DNA can be detected in the tumour cells by in-situ hybridization. Immunohistochemical expression of EBV latent membrane protein 1 is more variable [377,857,986,1316,1479,2326].

Differential diagnosis
Important differential diagnoses include metastatic undifferentiated carcinoma, malignant lymphoma, lymphoepithelial sialadenitis (no definite cytological atypia, presence of basement membrane-like material, no desmoplastic stroma, no EBV association), lymphadenoma (definite or subtle gland formation, no definite cytological atypia, no desmoplastic stroma, and no EBV association), and large cell undifferentiated carcinoma.

Precursor lesions
Most LEC arise de novo but rarely they may develop within lymphoepithelial sialadenitis (formerly myoepithelial sialadenitis) [938].

Genetic susceptibility
Clustering of salivary gland LEC in family members has been reported [31,91,1708]. One such family also showed dominantly inherited trichoepitheliomas, suggesting hereditary predisposition related to tumour suppressor genes [1708].

Prognosis and predictive factors
Five-year survival rate of 75-86% has been reported in patients treated by combined surgery (including neck dissection) and radiation therapy, although local recurrence can occur [236,1387, 1479,2252,2636]. The prognosis is significantly related to tumour stage. There have been attempts to grade LEC based on nuclear pleomorphism and mitotic activity [459,1373], with suggestion that high-grade tumours are more aggressive, but there are currently no widely accepted or well-validated grading systems.

Fig. 5.45 Lymphoepithelial carcinoma of parotid gland. A Carcinoma cells are admixed with many small lymphocytes. Note indistinct cell borders, vesicular nuclei and prominent nucleoli. B Tumour islands are heavily infiltrated by lymphocytes. C Uncommon cystic change in the tumour islands.

Fig. 5.46 Lymphoepithelial carcinoma of salivary gland. A Immunostaining for cytokeratin highlights the irregular tumour islands. B In-situ hybridization for EBV (EBER) selectively highlights the islands of carcinoma. The lymphoid cells in the background are negative.
Sialoblastoma

**Definition**
This is a rare, potentially aggressive, parotid or submandibular tumour that is usually present at birth and recapitulates the primitive salivary anlage.

**ICD-O code**
8974/1

**Synonyms**
Congenital basal cell adenoma, basal cell adenoma, basaloid adenocarcinoma, congenital hybrid basal cell adenoma-adenoid cystic carcinoma, embryoma (2570, 2685).

**Epidemiology**
Most tumours are identified at birth or shortly thereafter; occasional children may be diagnosed after the age of two years. The male to female ratio is 2:1. Sialoblastomas are extremely rare; 23 such cases have been reported (48, 156, 251, 867, 945, 1016, 1574, 1688, 1786, 1952, 2353, 2570, 2685).

**Localization**
The ratio of parotid to submandibular gland involvement is approximately 3:1.

**Clinical features**
Most babies present with a mass of the cheek or submandibular region. Occasional tumours may reach massive proportions and ulcerate skin. One baby presented with a concomitant hepatoblastoma (2353), and two other children both had congenital nevi associated with their tumours (251, 945). Some babies have been diagnosed by prenatal sonography. Radiographically, these tumours appear as expansile, lobulated masses. True-cut preoperative biopsy can be diagnostic, and is useful in ruling out neoplasia that require neoadjuvant chemotherapy, such as rhabdomyosarcoma.

**Histopathology**
Sialoblastomas are composed of basaloid epithelial cells, with scanty cytoplasm, round to oval nuclei, single or few nucleoli, and relatively fine chromatin pattern. More mature cuboidal epithelial cells with pink cytoplasm can also be seen. These cells form ductules, bud-like structures and solid organoid nests, and may demonstrate peripheral palisading. The intervening stroma may appear loose and immature. Myoepithelial cells can be identified, and have been confirmed by ultrastructural study. More familiar salivary patterns such as adenoid cystic-like cribriform areas can be seen. The mitotic rate within sialoblastomas is highly variable, and may increase with subsequent recurrences (251), as may necrosis, nuclear pleomorphism and MIB1 proliferative index.

**Immunoprofile**
These tumours express S-100 and vimentin diffusely. Cytokeratin accentuates the ductal structures.

**Histogenesis**
It has been suggested that these tumours originate from retained blastemal cells rather than basal reserve cells (2570). Dysembryogenic parotid changes have been described adjacent to the tumour, with proliferation of the terminal ductal epithelial bulbs (1952).

**Prognosis and predictive factors**
Sialoblastomas have the potential to recur (22%), and can occasionally metastasize regionally (9%), and one fatality has been reported (251, 1688). Most of these children are cured by primary surgical resection.

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Fig. 5.47 Sialoblastoma. **A** Solid nests composed of basaloid cells. **B** Brisk mitotic rate within this sialoblastoma.
Pleomorphic adenoma

Definition

Pleomorphic adenoma is a tumour of variable capsulation characterized microscopically by architectural rather than cellular pleomorphism. Epithelial and modified myoepithelial elements intermingle most commonly with tissue of mucoid, myxoid or chondroid appearance.

Synonym

Mixed tumour

ICD-O code

8940/0

Epidemiology

Pleomorphic adenoma is the most common salivary gland tumour and accounts for about 60% of all salivary neoplasms (2439). The reported annual incidence is 2.4-3.05 per 100,000 population (244, 2053). The mean age at presentation is 46 years but the age ranges from the first to the tenth decades (703). There is a slight female predominance (703,2711).

Localization

About 80% of pleomorphic adenomas arise in the parotid, 10% in the submandibular gland and 10% in the minor salivary glands of the oral cavity, nasal cavity and paranasal sinuses and the upper respiratory and alimentary tracts (703). The lower pole of the parotid gland is the most common location but deep lobe tumours can present as a parapharyngeal mass. The accessory parotid is occasionally involved.

Clinical features

Pleomorphic adenomas usually are slow growing painless masses. Small tumours typically form smooth, mobile, firm lumps but larger tumours tend to become bossellated and may attenuate the overlying skin or mucosa. Multifocal, recurrent tumours may form a fixed mass. Pleomorphic adenomas are usually solitary but they may show synchronous or metachronous association with other tumours, particularly Warthin tumour, in the same or other glands (2298). Pain or facial palsy are uncommon but are occasionally seen, usually in relation to infarcted tumours. The size of most tumours varies from about 2-5 cm but some reported cases have been massive (388). In the palate, tumours are usually seen at the junction of the hard and soft palate unilaterally. In the hard palate they feel fixed due to the proximity of the underlying mucoperiosteum.

Histopathology

Pleomorphic adenoma shows a remarkable degree of morphological diversity. The essential components are the capsule, epithelial and myoepithelial cells, and mesenchymal or stromal elements. The capsule varies in thickness and presence. A quantitative study showed the thickness ranged from 15-1750 mm (2732). When tumours were serially sectioned areas of capsular deficiency were seen in all cases (1418). In predominantly mucoid pleomorphic adenomas, the capsule may be virtually absent and the tumour abuts onto the adjacent salivary gland. Most tumours show areas where finger-like processes extend into the capsule. In addition, the tumour sometimes bulges through the capsule and forms what appear to be separate satellite nodules. These satellites are invariably attached to the main tumour by an isthmus (1418,1986). There is a tenden-
cy for clefts to form close to and parallel with the capsule. These clefts are within the tumour itself and leave tumour cells attached to the capsular wall. The epithelial component shows a wide variety of cell types including cuboidal, basaloid, squamous, spindle cell, plasmacytoid and clear cells. Rarely, mucous, sebaceous and serous acinar cells are seen. These cells are cytologically bland and typically have vacuolated nuclei, without prominent nucleoli, and a low mitotic frequency. The epithelium usually forms sheets or duct-like structures. There is a wide range of epithelial cellularity; sometimes, the epithelial component forms the bulk of the tumour (cellular pleomorphic adenoma). This phenomenon has no prognostic significance. The ducts show cuboidal luminal cells and there may be an abluminal layer of myoepithelial cells. These may be morphologically similar to the luminal cells or have clear cytoplasm and somewhat angulated nuclei. In limited material tumours showing these features could easily be confused with adenoid cystic carcinoma and epithelial-myoepithelial carcinoma. The ducts often contain eosinophilic secretory material and are usually small but may be distended to form microcysts. Squamous metaplasia, sometimes with the formation of keratin pearls, can be seen in both ducts and sheets and occasionally there is mucous metaplasia or conspicuous clear cell change. These appearances can be confused with mucoepidermoid carcinoma. More rarely, sebaceous cells or serous cells with zymogen granules are seen. Another rare feature is the presence of multinucleated epithelial cells. Myoepithelial cells may form a fine reticular pattern or sheets of spindle-shaped cells. These may be palissaded forming a Schwannoma-like appearance. A very distinctive appearance is seen when the myoepithelial cells are plasmacytoid or hyaline (1552). Focal oncocytyic change is not uncommon but occasionally the entire tumour is affected and may be mis-diagnosed as an oncocytoma (1973). Crystalloid material in the form of collagenous crystalloids, tyrosine and oxalate crystals are occasionally present (324).

The mesenchymal-like component is mucoid/myxoid, cartilaginous or hyalinised and sometimes this tissue forms the bulk of the tumour. Cells within the mucoid material are myoepithelial in origin and their cellular periphery tends to blend into the surrounding stroma. The cartilage-like material appears to be true cartilage and is positive for type II collagen and keratan sulphate. Occasionally it is the major component of the tumour. Bone may form within this cartilage or form directly by osseous metaplasia of the stroma. Deposition of homogeneous, eosinophilic, hyaline material between tumour cells and within the stroma can be a striking feature of some tumours. It forms globular masses or sheets and

Fig. 5.50 Pleomorphic adenoma. A Epithelial component with ductal structures (left) and a mesenchymal myxoid component (right). B Ducts showing luminal cells and several layers of abluminal cells, the latter being merged into myxoid stroma. C Cellular type. D Plasmacytoid cells.
Tumours of the salivary glands

typically is positive with stains for elastin. This material can push apart epithelial elements to give a cylindromatous or cribriform appearance that is readily mistaken for adenoid cystic carcinoma. Some longstanding tumours show increasing hyalinisation and the epithelial component is progressively effaced. It is important, however, to scrutinise the residual epithelial elements of such old, scarred pleomorphic adenomas as there is a significant risk of malignant progression in such tumours [85]. Tumours that have a lipomatous stromal component of 90% or more have been called lipomatous pleomorphic adenomas [1881, 2299].

More extensive inflammation and necrosis can be seen following spontaneous infarction or fine needle aspiration. In such tumours there may be an increase in mitotic figures and some cellular atypia [361,1495]. In addition, squamous metaplasia may be present and these changes can be mistaken for malignancy. Some tumours show cystic degeneration with the neoplastic elements forming a rim around a central cavity. Occasionally tumour cells can be seen within vascular spaces [475]. These are usually within the body of the tumour or at the periphery and this is assumed to be a peroperative phenomenon. Sometimes this is seen in vessels distant from the main tumour mass. However, this finding does not appear to have any significance in terms of tumour behaviour and, in particular, the risk of metastasis.

Immunoprofile

The inner ductal cells in the tubulo-glandular structures are positive for cytokeratin 3, 6, 10, 11, 13, and 16, whereas the neoplastic myoepithelial cells are irregularly positive for cytokeratin 13, 16, and 14 [311]. The neoplastic myoepithelial cells co-express vimentin and pan-cytokeratin and are variably positive for S-100 protein, a-smooth muscle actin, GFAP, calponin, CD10 and muscle-specific actin (HHF-35) [545]. Modified myoepithelial cells in these tumours are also reactive for p63 [214]. The non-lacunar cells in the chondroid areas are positive for both vimentin and pan-cytokeratin, whereas the lacunar cells are positive only for vimentin [1776]. The spindle-shaped neoplastic myoepithelial cells around the chondroid areas express bone morphogenetic protein (BMP) [1083] whereas the inner ductal cells in the tubulo-glandular structures and the lacuna cell in the chondroid areas express BMP-6 [1397]. Type II collagen and chondromodulin-I is present in the chondroid matrix [1396]. Aggrecan is present not only in the chondroid matrix but also in the myxoid stroma and in the inter-cellular spaces of the tubulo-glandular structures [2898].

Genetics

cytogenetics

Extensive cytogenetic studies of pleomorphic adenomas have shown that approximately 70% of the tumours are...
karyotypically abnormal [306,1639, 2239]. Four major cytogenetic subgroups may be discerned:

> Tumours with rearrangements involving 8q12 (39%)
> Tumours with rearrangements of 12q13-15 (8%)
> Tumours with sporadic, clonal changes not involving 8q12 or 12q13-15 (23%)
> Tumours with an apparently normal karyotype (30%).

Whereas t(3;8)(p21;q12) and t(5;8)(p13;q12) are the most frequently observed translocations in the first subgroup, a t(9;12)(p24;q14-15) or an ins(9;12)(p24;q12q15) are the most frequent rearrangements seen in the second subgroup. In addition, many variant translocations have been identified in which a number of other chromosome segments are found as translocation partners of both 8q12 and 12q13-15. Secondary chromosome changes, including trisomies, dicentrics, rings and double minutes, are found in about one-third of the cases with abnormal karyotypes. Previous studies have also indicated that patients with karyotypically normal adenomas are significantly older than those with rearrangements of 8q12 (51.1 years versus 39.3 years, p < 0.001) and that adenomas with normal karyotypes are often more stroma rich than tumours with 8q12 abnormalities [306].

**Molecular genetics**

The target gene in pleomorphic adenomas with 8q12 abnormalities is PLAG1, a developmentally regulated zinc finger gene [82,1279,2701]. Translocations involving 8q12 commonly result in promoter swapping/substitution between PLAG1 and a ubiquitously expressed translocation partner gene, leading to activation of PLAG1 expression. The breakpoints invariably occur in the 5’-noncoding regions of both the target gene and the promoter donor genes. The most commonly observed fusions are CTNNB1-PLAG1 and LIFR-PLAG1, resulting from t(3;8)(p21;q12) and t(5;8)(p13;q12) translocations, respectively [1279,2701]. Recently, cryptic gene fusions involving CTNNB1-PLAG1 and SII-PLAG1 were also found in karyotypically normal adenomas [82]. The PLAG1 protein is a nuclear oncoprotein that functions as a DNA-binding transcription factor. Deregulation of PLAG1 target genes, including IGF2, is likely to
play a major role in the genesis of pleomorphic adenomas [2700].

The target gene in adenomas with rearrangements of 12q14-15 is the high mobility group protein gene, HMGA2 (a.k.a. HMGIC) [878,879,2269]. HMGA2 encodes an architectural transcription factor that promotes activation of gene expression by modulating the conformation of DNA. The protein contains three DNA-binding domains that bind to the minor groove of AT-rich DNA. The majority of breakpoints in HMGA2 occur within the third large intron, resulting in separation of the DNA-binding domains from the highly acidic, carboxy-terminal domain. Two fusion genes, HMGA2-NFIB and HMGA2-FHIT, have been identified in adenomas with ins(9;12) and t(3;12), respectively [878,879]. Since no common functional domain has been found among the translocation partners, the critical event seems to be the separation of the DNA-binding domains from potential mRNA destabilizing motifs in the 3′-UTR, leading to deregulation of HMGA2 oncoprotein expression. High-level expression of HMGA2 resulting from gene amplification was recently suggested to be of importance for malignant transformation of pleomorphic adenomas [2194].

The five PLAG1- and HMGA2-containing fusion genes so far identified are all tumour specific and may therefore be used as diagnostic markers for pleomorphic adenomas [2194].

In tumours with PLAG1 activation [1727, 2198,2464,2465], whereas amplification and/or overexpression of ERBB2 seem to be rare [2198,2465]. Similarly, TP53 alterations are infrequent in adenomas [1907, 2198,2734]. In contrast, mutation and overexpression of TP53 are found in a relatively high proportion of carcinoma ex pleomorphic adenomas [1491,1907, 2169]. In addition, recent studies have shown that the TP53-related genes TP63 and TP73, which are novel myoepithelial markers, are overexpressed in basal and myoepithelial cells in pleomorphic adenomas [214,2734]. The pathogenetic relevance of the latter observations is uncertain. Studies using the human androgen receptor gene assay have demonstrated that the stromal and epithelial cells in pleomorphic adenomas are clonal and derived from the same progenitor cell [1455].

Finally, it was recently demonstrated that pleomorphic adenomas contain Simian virus 40 (SV40) DNA sequences and express the SV40 large T antigen, suggesting that this oncogenic virus may be involved in the genesis and/or progression of this tumour [1643].

Prognosis and predictive factors

Although pleomorphic adenoma is a benign tumour it can cause problems in clinical management due to its tendency to recur and the risk of malignant transformation. Recurrences are rare in the minor glands but in a meta-analysis of parotid tumours 3.4% of tumours recurred after 5 years and 6.8% after 10 years with a range of 1-50% [1083]. The variation of frequency of recurrence in this survey probably reflected the inclusion of cases reported before superficial parotidectomy became a widely used treatment and the variability of long-term follow-up. Some single centre, long-term surveys however, have shown recurrence rates as low as 1.6% [2169]. Recurrences appear to be much more likely in younger patients [1436,1681]. The possible reasons for recurrences or persistence in pleomorphic adenoma include:

- The diffusent nature of predominantly mucoid tumours [2157].
- The variability of the thickness of the capsule, together with the tendency of the tumour to invade the capsule [1085].
- Tumour nodules bulging through the capsule.
- Intratumoural splitting beneath the capsule.
- It is probable that the tumour cells have low biological requirements and this enables them to survive when spilt into the operative site.

Many recurrent pleomorphic adenomas are multifocal and some are so widely distributed that surgical control becomes impossible.

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**Fig. 5.55** Pleomorphic adenoma. A Schematic illustration of promoter swapping between PLAG1 and CTNNB1 in pleomorphic adenomas with t(3;8). Coding exons are indicated by filled boxes. Breakpoints are indicated by arrows. B Schematic illustration of the HMGA2 gene. Coding exons are indicated by filled boxes. Breakpoints are indicated by arrows.
Myoepithelioma

Definition
Myoepithelioma is a benign salivary gland tumour composed almost exclusively of sheets, islands or cords of cells with myoepithelial differentiation that may exhibit spindle, plasmacytoid, epithelioid or clear cytoplasmic features.

ICD-O code 8982/0

Synonyms
Myoepithelial adenoma, benign myoepithelial tumour.

Epidemiology
Myoepitheliomas account for 1.5% of all tumours in the major and minor salivary glands and represent 2.2% and 5.7%, respectively of all benign major and minor salivary gland tumours [668]. Both sexes are affected with equal frequency [41,128,546,668,1647,2282,2367]. Most tumours occur in adults, but rare examples have been recorded in children [1527]. The age of patients with myoepithelioma ranges from 9-85, with an average of 44 years and the peak age of occurrence in the third decade [668].

Localization
Myoepitheliomas develop preferentially in the parotid gland (40%) [668]. Minor salivary glands follow in frequency, especially in hard and soft palates [546,668,2282,2367]. Other minor salivary gland sites can also be affected [41,1647].

Clinical features
Myoepitheliomas usually present as slow growing painless masses [41,2282,2367].

Macroscopy
Myoepitheliomas are well-circumscribed, solid tumours that usually measure less than 3 cm in diameter [41,541,2367]. Myoepitheliomas have a solid, tan or yellow-tan, glistening cut surface [668].

Histopathology
A variety of cell morphologies has been recognized, including spindle, plasmacytoid or hyaline, epithelioid, and clear [546]. Most are composed of a single cell type but combinations may occur. Spindle cells are arranged in interlacing fascicles with stroma-like appearance [1579]. Plasmacytoid cells are polygonal cells with eccentric nuclei and dense, nongranular or hyaline, abundant eosinophilic cytoplasm. Plasmacytoid cells are found more often in tumours arising in the minor salivary glands than in the parotid gland. These hyaline cells may simulate neoplastic plasma cells, skeletal muscle or “rhabdoid” cells [1575]. Epithelioid cells are arranged in nests or cords of round to polygonal cells, with centrally located nuclei and a variable amount of eosinophilic cytoplasm. The surrounding stroma may be either collagenous or mucoid. Some myoepitheliomas are composed predominantly of clear polygonal cells with abundant and optically clear cytoplasm, containing large amounts of glycogen but devoid of mucin or fat. These tumours may show intercellular microcystic spaces. In other myoepitheliomas, occasional duct-like structures and intercellular microcystic spaces may be present. An unusual reticular variant of myoepithelioma characterized by netlike arrangements of interconnected cell cords, extending through a loose, vascularized stroma, has been reported [546].

Immunoprofile
The cells of myoepithelioma are usually positive for cytokeratins, especially for CK7 and 14. The reactivity of the spindle cells is variable for α-smooth muscle actin, muscle specific actin (MSA), calponin, S-100, GFAP and smooth muscle myosin heavy chain. There is considerable variation of tumour expression of MSA. The spindle cells react strongly for MSA, the epithelioid cells react sporadically, and the plasmacytoid and clear cells are often nonreactive [805].

Electron microscopy
Ultrastructural studies confirmed the epithelial and myoepithelial differentiation of myoepithelioma [538,541].

Differential diagnoses
Distinction from pleomorphic adenoma is based on the relative lack of ducts and the absence of myxochondroid or chondroid areas. Myoepitheliomas with clear cells, or mixed epithelioid and clear cells have to be separated from other salivary gland tumours with clear cells, such as: mucoepidermoid carcinoma, acinic cell carcinoma, epithelial-myoepithelial carcinoma, oncocytoma and clear cell carcinoma. All these tumours lack the characteristic immunoprofile of the myoepithelial cells. In contrast to carcinomas, myoepitheliomas have a non-infiltrative, well-circumscribed periphery.

Table 5.03 Classification of clear cell tumours of the salivary glands.

<table>
<thead>
<tr>
<th>Classification</th>
<th>Examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benign</td>
<td>Pleomorphic adenoma, myoepithelioma, sebaceous adenoma, oncocytoma and oncocytic hyperplasia</td>
</tr>
<tr>
<td>Malignant, primary</td>
<td>a) Carcinomas not usually characterized by clear cells, but with clear cell predominant areas; e.g. mucoepidermoid and acinic cell carcinomas.</td>
</tr>
<tr>
<td></td>
<td>b) Carcinomas usually characterized by clear cells;</td>
</tr>
<tr>
<td></td>
<td>i. Dimorphic epithelial-myoepithelial carcinoma.</td>
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<tr>
<td></td>
<td>ii. Monomorphic clear cell carcinoma.</td>
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<tr>
<td></td>
<td>iii. Sebaceous carcinoma.</td>
</tr>
<tr>
<td>Malignant, metastatic</td>
<td>Carcinomas, especially kidney, thyroid, melanoma.</td>
</tr>
</tbody>
</table>

A. Cardesa
L. Alos
Predominantly spindle cell myoepitheliomas must be distinguished from benign and malignant mesenchymal tumours.

**Genetics**
Cytogenetic studies have demonstrated structural alterations of chromosomes 1, 9, 12, and 13: t(1;12)(q25;q12), del(9)(q22.1q22.3), del(13)(q12q22) in a parotid myoepithelioma [654]. Mutations of TP53 have been observed in 3 of 12 (25%) myoepitheliomas [2734].

**Prognosis and predictive factors**
According to well-documented series myoepitheliomas are less prone to recur than pleomorphic adenomas [2282]. However, higher recurrence rates have been reported by others [41,646]. Recurrence is correlated with positive margins at the first excision [646]. The recommended treatment is complete surgical excision. Benign myoepitheliomas can undergo malignant transformation, especially in long standing tumours or in tumours with multiple recurrences [41].
**Basal cell adenoma**

**Definition**
Basal cell adenoma (BCA) is a rare benign neoplasm characterized by the basaloid appearance of the tumour cells and absence of the myxochondroid stromal component present in pleomorphic adenoma.

**ICD-O code**
8147/0

**Epidemiology**
Accurate epidemiological data are hard to obtain since in the past BCA was included within non-pleomorphic tumours. The BCAs are rare, accounting for 1-3% of all salivary gland tumours. They are typically seen in adults in the 7th decade with a 2:1 female predilection (2303), except for the membranous type that has an equal female:male distribution (668).

**Localization**
The majority arise in the major glands, and the parotid is the most frequent site of occurrence (~75%), followed by the submandibular gland (~5%) (162,2881). It is extremely rare in minor salivary glands, the upper lip being the most common site, followed by the buccal mucosa (704,2711).

**Clinical features**
Most tumours are solitary, well-defined, movable nodules. They are usually firm but occasionally cystic. The membranous type (dermal analogue tumour) (153) may be multiple and co-exist with dermal cylindromas or trichoepitheliomas (1033,1582,2867).

**Macroscopy**
Most of the tumours present as small, well-circumscribed, encapsulated nodules measuring between 1-3 cm, except for the membranous type that may be multinodular or multifocal. On cut section they are solid and homogeneous or cystic, with a greyish-white to brown colour.

**Histopathology**
Microscopically, BCAs are composed of basaloid cells with eosinophilic cytoplasm, indistinct cell borders and round to oval nuclei, distributed in solid, trabecular, tubular, and membranous patterns. However, tumours may present with more than one of these patterns, usually with the predominance of one. The solid type is composed of sheets or islands of variable shapes and sizes, usually with peripheral palisading of cuboidal to columnar cells. The islands are separated by strands of dense collagenous tissue. The trabecular type is characterized by narrow strands, trabeculae or cords of basaloid cells separated by cellular and vascular stroma. A rare but distinctive feature is the presence of a richly cellular stroma composed of modified myoepithelial cells (542). Ductal lumina are often observed among the basaloid cells and these cases are considered as tubulo-trabecular type. The membranous type of BCA has thick bands of hyaline material at the periphery of basaloid cells and as intercellular coalescing droplets. In the tubular type, ductal structures are a prominent feature. All variants may demonstrate cystic change, squamous differentiation in the form of whorls or ‘eddies’, or rare cribriform patterns. Occasional tumours, particularly of the tubular type, are largely oncocytic.

**Immunoprofile**
Immunopositivity for keratin, myogenic markers, vimentin and p63 indicate ductal and myoepithelial differentiation (214, 553,1598,2883). Also the palisading cells of the solid type can stain for vimentin and myogenic markers. The pattern of expression reflects the different differentiation stages of the tumour cells, varying from the solid type, the less differentiated, to the tubular type, the most differentiated.

**Genetics**
Genetic aberration has been described in three cases of BCA. Two cases presented trisomy 8 and one case the 7;13 translocation and/or inv(13) (1136,2385).

**Prognostic and predictive factors**
BCA is usually a non-recurrent tumour, except for the membranous type, that has a recurrence rate of approximately 25% (1582). Although exceedingly rare, malignant transformation of BCA has been reported (1825).
Fig. 5.59 Histological types of basal cell adenoma. A Tubular type, with small duct lumens lined by cuboidal eosinophilic cells. B Membranous type, with prominent hyaline material around and inside epithelial islands. C Occasional features found in basal cell adenoma include variable sized cystic spaces. D High cellularity of the stroma represented by spindle-shape cells.

Fig. 5.60 Immunohistochemical profile of basal cell adenoma. Tubulo-trabecular type. A CK7 positivity in ductal cells. B Smooth muscle actin expression in myoepithelial cells.
Warthin tumour

Definition
A tumour composed of glandular and often cystic structures, sometimes with a papillary cystic arrangement, lined by characteristic bilayered epithelium, comprising inner columnar eosinophilic or oncocytic cells surrounded by smaller basal cells. The stroma contains a variable amount of lymphoid tissue with germinal centres.

ICD-O code 8561/0

Synonyms
Adenolymphoma, cystadenolymphoma, papillary cystadenoma lymphomatosum. Warthin tumour is preferred to avoid any possible confusion with a lymphoid malignancy, and with the separate entity, lymphadenoma [1591].

Epidemiology
In most countries, Warthin tumour is the second commonest tumour of the salivary glands. In the United States (US) it comprised about 3.5% of all primary epithelial tumours (5.3% in the parotid) [668]. Other studies revealed higher percentages, such as 14.4% of primary epithelial tumours of the parotid gland in the United Kingdom (UK) [703], 27% in Denmark [2075], and 30% in Pennsylvania, USA [1765]. Warthin tumour occurs in Caucasians and Asians [451], but has a lower incidence in African-Americans [668] (although this may now be increasing [2856]) and in Black Africans [2590]. The mean age at diagnosis is 62 years, (range 12-92) [668], and it is rare before 40. The relative sex incidence has changed during the last half-century: In 1953 the male to female ratio was 10:1 [786], whereas in 1996 it was 1.2:1 [668], and in 1992 it was equal [1765]. In the UK in 1986 the ratio was 1.6:1 [705].

Localization
Warthin tumour is almost exclusively restricted to the parotid glands and the periparotid lymph nodes. Most cases involve the lower pole although 10% are in the deep lobe. Occasional tumours (2.7% in one series) arise within adjacent lymph nodes [664]. Very rare examples have been reported in other glands [2669], but some tumours thought initially to be within the submandibular gland have usually arisen from the anterior tail of the parotid or from lymph nodes [668]. Warthin tumour is clinically multicentric in 12-20% of patients (either synchronous or metachronous), and is bilateral in 5-14% [899,1610]. In addition, serial sectioning revealed additional sub-clinical lesions in 50% of cases [1417].

Etiology
There is a strong link between Warthin tumour and cigarette smoking [633,2052] – the incidence is eight times that of non-smokers [1360]. In addition, the increased numbers of female smokers during the second half of the 20th century closely parallels the increase in Warthin tumour in women, and largely explains the change in sex incidence during this period [1421,2856]. The mechanisms are not clear but in has been speculated that irritants in tobacco smoke cause metaplasia in the parotid [2866]. Radiation exposure may be relevant as there is an increase in Warthin tumour among atomic bomb survivors [2229]. There is also said to be a higher frequency of autoimmune disorders in patients with Warthin tumour than in those with pleomorphic adenomas or healthy subjects [899]. At present, the balance of probabilities is that EBV does not play a significant role in the etiology of Warthin tumour [2733]. The metaplastic (infarcted) variant can follow trauma, particularly from FNA biopsy [596,706].

Clinical features
Most patients present with a painless mass, on average, 2-4 cm, although...
occasional cases have reached 12 cm (2783). The mean duration of symptoms is 21 months, but in 41% of patients it is less than six months (705). Many patients notice fluctuation in size of the tumour, especially when eating (1711). Pain has been reported in 9% (705), particularly those with the metaplastic variant (2866). Facial paralysis is very rare, and is the result of secondary inflammation and fibrosis, and likewise can be seen in the metaplastic variant (706,1876).

Warthin tumour is able to concentrate Technetium (99mTc), appearing as a “hot” lesion. It is usually well-circumscribed, but secondary inflammation can cause the edges to become indistinct.

**Macroscopy**
Most Warthin tumours are well-circumscribed, spherical to ovoid masses, and partly cystic. The cysts vary from small slits to spaces up to several centimetres, and contain clear, mucoid, creamy white or brown fluid. Solid areas are tan to white, and often firm and fibrous in the metaplastic variant. In all cases of Warthin tumour, the parotidectomy specimen should be examined for other lesions.

**Histopathology**
The tumour is sharply demarcated with a thin capsule. There are cystic and solid areas, composed of epithelial and lymphoid components. The cysts and slit-like spaces vary in size and shape, and papillary structures project into the lumen. The papillae have fibrovascular cores often with lymphoid stroma. The epithelium comprises two layers of cells: the oncocytic luminal cells are tall and columnar, and show palisading of their bland single ovoid nuclei. The surface often shows apocrine blebbing and cilia are occasionally identified (705). Deep to this layer lie smaller flattened or cuboidal basal cells. Their cytoplasm is similar, but less abundant. No significant nuclear atypia or mitotic activity is identified. Small foci of squamous metaplasia, scanty goblet cells and very occasional sebaceous cells are seen.

The stroma comprises lymphoid tissue displaying varying degrees of reactivity, and germinal centres are usual. Increased numbers of mast cells and plasma cells may also be seen. The cystic spaces contain eosinophilic secretions with occasional crystal formation and laminated bodies resembling corpora amylacea.

Some tumours, variously termed, infarcted, infected or metaplastic, account for 6-7% of Warthin tumours (706,2295). They are likely to be encountered more frequently in the future with the increasing use of pre-operative FNA. There is extensive necrosis, in which a ghost architecture of papillary structures is often identified – this can be highlighted with a reticulin stain. Non-keratinizing squamous metaplasia is prominent, consisting of tongues and cords of often spongiotic squamous cells extending into surrounding tissues in a pseudo-infiltrative pattern. Cytological atypia can be prominent, and mitotic figures numerous, but none is abnormal. Goblet cells can also be seen, but should not be numerous. At the periphery of the lesions, there is extensive fibrosis, with dense hypocellular collagen and myofibroblastic spindle cell proliferation. There is a heavy mixed inflammatory infiltrate, comprising neutrophils, chronic inflammatory cells, as well as sheets of macrophages, some with foamy cytoplasm. Lipogranulomas, with or without cholesterol crystals, are not uncommon. Areas of residual undamaged Warthin tumour can be found, but not in every case, and there may thus be few clues to the nature of the original lesion (596,706).

**Immunoprofile**
Lymphoid marker studies have shown B (CD20), NK (CD56) and T (CD3) cells, including helper (CD4) and suppressor (CD8) subtypes. This profile of lymphocyte subsets is similar to that in normal or reactive lymph nodes (432). Special stains and immunohistochemistry have little to offer in the diagnosis of Warthin tumour, although there may be a role in diagnosing the metaplastic variant with epithelial markers, particularly when no residual viable Warthin tumour can be identified (596,2279).

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Fig. 5.62 Warthin tumour. A Conspicuous mucous metaplasia. B Luminal layer of palisaded oncocytic cells with apocrine blebs, and less numerous abluminal cells.

Fig. 5.63 Warthin tumour. Electron micrograph showing ciliated epithelium.
Cytopathology
The cytopathological findings reflect the histopathological appearance, except that mast cells are more noticeable. The other cellular elements are oncotypic epithelial cells and lymphocytes, with a background of cell debris and proteinaceous material [776]. Uncommon findings include ciliated cells [2899], squamous cells, mucous cells, siderophages, giant cells, calcifications and crystalloids [776]. The diagnostic sensitivity of FNA cytology is moderately accurate [1984], but the error rate is clinically significant, as for example, the findings in lymphocyte-rich acinic cell carcinoma are almost identical [2135].

Differential diagnosis
Of all salivary gland tumours, the typical type of Warthin tumour is usually unmistakable. Papillary cystadenoma is similar and possibly related, but any lymphoid tissue is scanty. There is some resemblance to other lymphoepithelial cystic lesions such as simple benign lymphoepithelial cyst (unrelated to AIDS), lymphoepithelial sialadenitis (LESA) with cystically dilated ducts, cystic lymphoid hyperplasia of AIDS and MALT lymphoma with cystically dilated ducts [2372]. An important differential is from cystic metastases in intra and parotid lymph nodes – the malignant nature of most should be obvious, but a recently-reported variant of papillary thyroid carcinoma has been described as “Warthin-like” [113,1572]. It is characterised by a heavy lymphoid stroma and oncotypic metaplasia of the epithelium. The best guide to its true nature is that the nuclei display typical chromatin clearing, inclusions and groove-formation, and the epithelial cells show immunohistochemical expression of thyroglobulin. If there is marked cytological atypia and mitotic activity, the metaplastic variant can be mistaken for squamous or mucoepidermoid carcinoma, either primary or metastatic [596]. The resemblance is particularly close if there has been total infarction of the original Warthin tumour. Clues to the true nature of the lesion include any ghost papillary architecture in the necrotic zones. Also, the squamous metaplasia lacks keratinization (seen in most squamous carcinomas), and mucinous goblet cells are usually much less numerous than in low-grade cystic mucoepidermoid carcinoma.

Histogenesis
There are two principal theories of the histogenesis [668]: one is an origin from intercalated and basal cells of heterotopic salivary ductal inclusions in intra- or peri-parotid lymph nodes. In particular, this explains the distribution of Warthin tumour and its absence from other salivary tissue lacking incorporated lymph nodes. The alternative theory is that Warthin tumour is a benign epithelial neoplasm or proliferation that attracts a heavy lymphoid reaction, similar to that seen in certain other salivary neoplasms [83,1717,2372]. More recently, it has been suggested that Warthin tumour initially develops in a parotid lymph node as an adenomatous epithelial proliferation responding to as yet unidentified stimuli (probably including tobacco either as a direct stimulus or a promoter), followed by lymphocytic infiltration. The stage of this process seen at the time of surgery determines the proportions of epithelial and lymphoid elements [20].

Genetics
Cytogenetic studies have shown Warthin tumour to have three main stemline groups, one with a normal karyotype, a second with numerical changes only (loss of Y chromosome or trisomy or monosomy 5) and a third group involving structural changes with one or two reciprocal translocations [1711]. Damage to the mitochondrial DNA may account for the ultrastructural changes seen in the mitochondria, as well as the oncotypic change seen morphologically [1494]. Analysis of the X chromosome-linked human androgen receptor gene showed that Warthin tumour is non-clonal, and thus likely to be non-neoplastic [1118]. This finding supports morphological observations that suggested Warthin tumour (as well as various thymic and head and neck cysts) resulted from the induction of cystic changes in branchial cleft epithelium by an inflammatory infiltrate, accompanied by oncotypic change in the epithelium [2199,2509]. A study of 13 cystadenolymphomas (Warthin tumours) showed minimal chromosomal alterations in these tumours [1905]. Interestingly, at least two tumours with cytogenetic analysis have been reported to have t(11;19) (q21;p13) translocation, suggesting a link to mucoepidermoid carcinoma [305,1638]. It is interesting, that the rearrangements on 8q and 12q have, so far, been found to be mutually exclusive [306,2239].

Prognosis and predictive factors
Primary treatment is surgical, either superficial parotidectomy or enucleation. After this, most studies show low recurrence rates of about 2-5.5% [668,705], presumably the result of multifocality. Malignant change is rare, at about 1% [669,2295], and may involve the epithelial or lymphoid components. Some patients give a history of radiation [1984,2229,2295]. Several types of carcinoma have been described, including squamous [2390], adenocarcinoma [2295], mucoepidermoid [1826,2294], oncotypic [2585], Merkel cell [788] and undifferentiated. The differential diagnosis includes squamous or mucous metaplasia, and metastases of extra-salivary malignancies to a pre-existing Warthin tumour. Lymphomas include nodal types [115,1694,2338], and one report of lymphoepithelial lesions suggesting a MALT-type neoplasm [113]. Warthin tumour is sometimes seen in association with other benign salivary tumours, particularly pleomorphic adenoma [664,905,1458,2338,2395], although it is not clear if this is greater than would be expected by chance with what is after all not an uncommon tumour. Another study found an increased incidence of extra-salivary neoplasms. A common etiology of cigarette smoking explains the carcinomas of the lung, larynx and possibly the bladder, whilst the others (lymphoma, kidney and breast cancers) could just be a coincidence [1610].
**Definition**
Benign tumour of salivary gland origin composed exclusively of large epithelial cells with characteristic bright eosinophilic granular cytoplasm (oncocytic cells).

**ICD-O code**
8290/0

**Synonym**
Oncocytic adenoma, oxyphilic adenoma

**Epidemiology**
Oncocytoma accounts for about 1% of all salivary gland neoplasms and occurs most commonly in the 6-8th decades (257). The mean age of the patients is 58 years. There is no sex predilection.

**Etiology**
Approximately 20% of all the patients will have a history of radiation therapy to the face or upper torso or long-term occupational radiation exposure five or more years prior to tumour discovery (257). Patients with previous radiation exposure are on the average 20 years younger at tumour discovery than those without a documented history of irradiation.

**Localization**
Among oncocytic major salivary gland tumours, 84% occur in the parotid (male to female ratio of 1:1), and the remainder arise in the submandibular gland (2601). Minor salivary gland sites include the lower lip, palate, pharynx, and buccal mucosa.

**Clinical features**
Symptoms vary according to the site of occurrence and most commonly present as a painless mass, less frequently nasal or airway obstruction.

**Imaging**
CT scan: Well-defined area of increased density in the host salivary gland. Radiouclide imaging shows increased uptake of technetium-99m that does not disappear following sialogogue administration. This finding plays an important role in the diagnosis and is related to the presence of oncocytes and their increased mitochondrial content.

**Macroscopy**
On gross examination, oncocytomas are usually 3-4 cm in size and possess a well-defined capsule. The cut surface is light brown and lobular.

**Histopathology**
Histologically, the oncocytes display ample granular acidophilic cytoplasm. Typically the predominant cells have abundant oncocytic cytoplasm and an oval, vesicular nucleus (light cells). In addition, there are cells with very brightly eosinophilic cytoplasm and pyknotic nuclei (dark cells). The cells are arranged in uniform sheets and they may aggregate into clusters, and sometimes they form duct-like structures. Rarely, oncocytomas present with large polyhedral clear cells in an organoid distribution. A thin fibrovascular stroma is also present. An intimate mixture of typical eosinophilic and clear cell oncocytes may be encountered within the same tumour. Tumours with a predominantly clear cell component are referred to as clear cell oncocytoma (665). The optical clear cell appearance is due to fixation artefact and/or intracytoplasmic glycogen deposition (551,2291). The tumour cells typically stain with phosphotungstic acid haematoxylin (PTAH). Electron microscopy shows elongated cristae and a partial lamellar internal structure (1227). The nuclei of the oncocytes are irregular and contain inclusions and glycogen granules.

**Differential diagnosis**
The most important differential diagnosis of oncocytoma includes acinic cell carcinoma and clear cell carcinoma. Mucoepidermoid carcinoma with prominent clear cell alteration and metastatic renal cell carcinoma may also be practical considerations. Also, stroma-poor Warthin tumour, oncocytic carcinoma, and metastatic thyroid carcinoma should be included. The clear-cut separation of an oncocytic adenomatous (nodular) hyperplasia of the parotid gland from a multinodular oncocytoma (a true neoplasm) is not always possible since the two entities overlap histologically (223,882,2427).

**Prognosis and predictive factors**
Complete surgical excision is the treatment of choice. Radiotherapy is not indicated especially since oncocytes are radioresistant. Local recurrence of an oncocytoma is extremely rare, but when it occurs, it may be multiple and bilateral. The incidence of bilateral oncocytomas is 7%. It seems there is an association between bilateral disease, tumour recurrence, and marked clear cell change (clear cell oncocytosis).
Canalicular adenoma

Definition
The tumour is composed of columnar epithelial cells arranged in thin, anastomosing cords often with a beaded pattern. The stroma is characteristically paucicellular and highly vascular.

ICD-O code 8149/0

Synonyms
Basal cell adenoma, canalicular type, monomorphic adenoma, canalicular type, adenomatosis of minor salivary glands

Epidemiology
There is a peak incidence in the seventh decade (mean 65 years). The age range is 33-87 years. It is uncommon before the age of 50 [529,668,1864] and the female-to-male ratio is 1.8:1 [529,668]. It comprised 1% of all salivary gland neoplasms and 4% of minor salivary gland neoplasms in a major series [668].

Localization
Canalicular adenoma has a peculiar predilection to involve the upper lip (about 80% of tumours) [529,1864]. The next most common location is the buccal mucosa (9.5% of tumours) [1864]. Rarely, canalicular adenoma can involve the major salivary glands [529].

Clinical features
These tumours present as enlarging nodules with no accompanying symptoms such as pain or paralysis. The overlying mucosa shows typical coloration but in some cases may appear bluish. A peculiar presentation of canalicular adenoma is that of multiple /multifocal canalicular adenomas [1308,1866,2206]. When this occurs, the upper lip and buccal mucosa are typically involved but other sites can be affected.

Macroscopy
Canalicular adenomas range in size from 0.5-2.0 cm in diameter and are grossly well circumscribed. The colour is light yellow to tan [668].

Histopathology
The microscopic appearance at low magnification likewise shows circumscript. Some canalicular adenomas have a fibrous capsule while smaller tumours often do not. It is not uncommon to see multifocal microscopic canalicular adenomas adjacent to a larger canalicular adenoma. In addition, very small foci of adenomatous tissue can be seen which may represent the earliest recognizable microscopic manifestation of canalicular adenoma. Superimposed necrosis can occur in some cases [36]. The epithelial component manifests as two rows of columnar cells which alternately are situated opposed to each other and alternately widely separated. This leads to the characteristic appearance of these tumours - canaliculi - where the epithelial cells are widely separated. The alternating arrangement of closely opposed and widely separated epithelial cells also leads to the characteristic beaded appearance of these tumours. The epithelial cells forming the cords are typically columnar but can be cuboidal. Nuclei are regular and show no pleomorphism. Nucleoli are inconspicuous and mitotic figures are rare. The stroma is characteristic and a useful clue to the diagnosis. It is paucicellular but shows a prominent vascular pattern. The capillaries often have an eosinophilic cuff of connective tissue.

Immunoprofile
Canalicular adenomas stain with anti-keratin, anti-vimentin and anti S-100 antibodies [758]. Rare focal GFAP positivity is seen [758]. Canalicular adenomas are devoid of staining when more sensitive markers of myogenous differentiation such as smooth muscle actin, smooth muscle myosin heavy chain and calponin are used [2883].

Differential diagnosis
The most important are adenoid cystic carcinoma and basal cell adenoma. Multifocality and cribriform pattern should not be misinterpreted as carcinoma. Hybrid tumours composed of canalicular adenoma and basal cell adenoma have been reported [2297].

Prognosis and predictive factors
The prognosis is excellent and recurrences are rare even if the tumours are treated with just a local excision or lumpectomy. Whether new tumours are true recurrences or are a manifestation of the multicentric growth pattern is difficult to ascertain.
Sebaceous adenoma

Definition
It is a rare, usually well-circumscribed tumour composed of irregularly sized and shaped nests of sebaceous cells without cytologic atypia, often with areas of squamous differentiation and cystic change.

ICD-O code 8410/0

Epidemiology
They account for 0.1% of all salivary gland neoplasms and slightly less than 0.5% of all salivary adenomas (2301). The mean age is 58 years (22-90 years) and the male:female ratio is 1.6:1 (896, 901). Unlike cutaneous sebaceous neoplasms (1132,2214), there is no increased risk of developing a visceral carcinoma.

Localization
Approximately 50% of tumours arise in the parotid gland, 17% in the buccal mucosa, 13% in the retromolar region or area of the lower molars and 8% in the submandibular region (896).

Clinical features
Patients typically present with a painless mass.

Macroscopy
These adenomas range in size from 0.4-3.0 cm in greatest dimension, are commonly well circumscribed to encapsulated and are greyish-white to yellow (896,901).

Histopathology
They are composed of sebaceous cell nests often with areas of squamous differentiation with minimal atypia and pleomorphism with no tendency to invade local structures. Many tumours are microcystic or composed predominantly of ectatic ductal structures. The sebaceous glands frequently vary markedly in size and tortuosity and are often embedded in a fibrous stroma. Occasional tumours demonstrate marked oncocytic metaplasia. Histiocytes and/or foreign body giant cells can be seen focally. Lymphoid follicles, cytologic atypia, cellular necrosis, and mitoses are usually not observed. Infrequently, these tumours may be part of a hybrid neoplasm (2297).

Treatment and prognosis
Treatment consists of complete surgical excision. They do not recur.
Lymphadenomas: sebaceous and non-sebaceous

Definition
Sebaceous lymphadenoma is a rare, well-circumscribed to encapsulated tumour composed of variably sized and shaped nests of sebaceous glands without atypia often intermixed with different proportions of variably sized ducts, within a background of lymphocytes and lymphoid follicles. Lymphadenoma is a similar tumour lacking sebaceous differentiation.

ICD-O code
Sebaceous lymphadenoma 8410/0

Epidemiology
Approximately 75% of sebaceous lymphadenomas are first diagnosed in the 6-8th decades of life (25-89 years). There is no sex predominance. Lymphadenoma is a rare tumour, with only 5 cases having been reported in the literature [83,1399,1591]. From the limited available data, all patients are male ranging in age from 17-57 years.

Localization
Well over 90% of sebaceous lymphadenomas occurred in or around the parotid gland with one tumour arising in the anterior midline of the neck [896], and two tumours occurring in the oral region [1393,1654]. All cases of lymphadenomas reported so far have occurred in the parotid gland [83,1591].

Clinical features
Patients typically present with a painless mass.

Macroscopy
Tumours have ranged from 1.3-6.0 cm in greatest dimension. They are usually encapsulated, solid, multicystic, or unicystic masses that range from yellow to grey. Sebum is commonly found in many of the cysts.

Histopathology
Sebaceous lymphadenoma.
The majority of sebaceous lymphadenomas are composed of variably sized sebaceous glands admixed with salivary ducts in a diffuse lymphoid background. Others consist mainly of lymphocytes and lymphoid follicles surrounding ductal structures with only occasional sebaceous glands. All tumours have a lymphoid background, and about one half have well-developed lymphoid follicles. In addition, tumours may contain small areas of identifiable residual lymph node and focal necrosis has rarely been observed. Occasional tumours may also contain or be intermixed with components of a Warthin tumour or membranous basal cell adenoma [896,901]. Histiocytes and foreign body giant cell inflammatory reactions secondary to extravasated sebum are commonly observed. This foreign body reaction can be helpful in differentiating these tumours from mucoepidermoid carcinoma (MEC). Unlike MEC, which contains a variety of cell types; mucin positivity is never found in the clear sebaceous cells. However, intracellular and extracellular mucin may be occasionally found within ducts adjacent to sebaceous cells.

Lymphadenoma
It can take the form of anastomosing trabeculae or solid tubules surrounded by basement membrane-like material, or cystically-dilated glands filled with proteinaceous materials. Papillary structures can be found in some cases. The lining cells are cuboidal to columnar and show no significant cytologic atypia or mitotic activity. Basal cells can be identified in some areas. However, the epithelial component can be obscured by abundant admixed and intraepithelial lymphocytes; the diastase-peroxidase acid Schiff stain can help in highlighting the basement membrane around the epithelial nests. The lymphoid stroma comprises dense populations of lymphoid cells with lymphoid follicle formation. The lymphoid component is generally considered to represent tumour-associated lymphoid proliferation [83,604], hence conventional salivary gland adenomas occurring within intraparotid or cervical lymph node are excluded.

Differential diagnosis
The most important differential diagnosis of lymphadenoma is lymphoepithelial carcinoma; distinguishing features of lymphadenoma are lack of mitotic activity, lack of invasive growth with desmoplastic stroma, presence of subtle or overt ductal differentiation, and absence of EBV association. Metastatic adenocarcinoma in lymph node is characterized by recognizable nodal structures, definite nuclear atypia and invasive growth. Lymphadenoma can be distinguished from lymphoepithelial sialadenitis by the circumscribed borders as well as a more proliferative epithelial component.

Treatment and prognosis
Treatment consists of complete surgical excision. These tumours rarely recur.
Ductal papillomas are a group of relatively rare, benign, papillary salivary gland tumours known as inverted ductal papilloma, intraductal papilloma, and sialadenoma papilliferum. They represent adenomas with unique papillary features with a common relationship to the excretory salivary duct system, a non-aggressive biologic behaviour, and a predilection for the minor salivary glands. They tend to occur in the middle-aged and elderly and rarely in children. The three types of ductal papillomas possess distinct clinical and histologic features allowing differentiation from each other and other adenomas with a papillary pattern.

**Inverted ductal papilloma**

**Definition**
Inverted ductal papilloma is a luminal papillary proliferation arising at the junction of a salivary gland duct and the oral mucosal surface epithelium and exhibits an endophytic growth pattern that forms a nodular mass.

**ICD-O code** 8503/0

**Synonym** Epidermoid papillary adenoma

**Epidemiology**
The true incidence of inverted ductal papilloma (IDP) is unknown, but it is thought to be relatively rare based on the sparse number of reported cases. Lesions have arisen in adults with an age range of 28-77 years and a male predilection [264].

**Localization**
All of the reported sites have been in the minor salivary glands—the most common location is the lower lip followed by the buccal mucosa/mandibular vestibule. Other reported sites have been the palate and the floor of mouth [264].

**Clinical features**
IDP typically presents as a painless nodular submucosal swelling, often with a dilated pore or punctum surfacing the swelling [1046]. Lesions have been described as being present from months to several years.

**Macroscopy**
Lesions have ranged from 0.5-1.5 cm. They are nodular masses that are often papillary and occasionally cystic.

**Histopathology**
The neoplasms are unencapsulated. Well demarcated endophytic epithelial masses that are typically continuous with the mucosal epithelium. The mucosal epithelium has a central pore-like opening in the mucosal surface. The peripheral borders of the epithelial mass show a broad, smooth “pushing” interface juxtaposed to the connective tissue stroma. The epithelium proliferates in broad papillary projections that extend into the luminal cavity and are composed predominantly of epidermoid and basal cells that show columnar epithelium on the surface of the papillae. Acinar aggregates or individual mucocytes can be found in the columnar epithelial layer and/or in the subjacent epidermoid component. The epithelial cells are cytologically bland with little or no pleomorphism. Mitotic figures are rare.

**Differential diagnosis**
IDP must be differentiated from mucoepidermoid carcinoma since both have epidermoid and mucous cells. Inverted ductal papilloma does not have the multicystic, multinodular, and infiltrative growth pattern of mucoepidermoid carcinoma. Papillary features are rarely found in mucoepidermoid carcinomas.

**Prognosis and predictive factors**
There have been no reported recur-
Intraductal papilloma

Definition
Intraductal papilloma is a luminal papillary proliferation of duct epithelium that arises from a segment of the interlobular or excretory duct and causes unicystic dilatation.

ICD-O code 8503/0

Epidemiology
The intraductal papilloma is very rare. Age range is 8-77 years with most cases occurring in the 6th and 7th decade of life (264,1375). Sex distribution is essentially even.

Localization
The minor salivary glands are more frequently involved than the major glands. Intraductal papillomas are most commonly found in the lips and buccal mucosa. Tumours have been reported in the palate and tongue as well. Of the major glands the parotid is most frequently involved, but cases in the submandibular and sublingual glands have also been cited (264,1008,1749).

Clinical features
Intraductal papillomas of major and minor salivary glands present as painless, well-defined solitary masses or swellings. Duration can range from weeks to years.

Macroscopy
Grossly, intraductal papillomas are well-circumscribed, unicystic nodules that range in size from 0.5-2.0 cm. The lumina contain finely granular, often friable tissue and mucinous material.

Histopathology
The tumour is entirely confined within a circumscribed or encapsulated unicystic cavity. The lumen is partially or completely filled with many branching papillary elements consisting of fibrovascular cores surfaced by columnar to cuboidal cells of one to two layers that originate from a focal point in the wall. Mucocytes, often goblet-like, are interspersed throughout the epithelium lining the papillary elements. These mucous-containing cells can be few to many in number. The epithelium that lines the cyst-like cavity is composed of the same type of epithelium as the papillary fronds. In many instances, the cystic structure has a dense fibrous connective tissue wall surrounding it. Cytologic atypia and mitotic figures are virtually absent (264).

Differential diagnosis
In contrast to intraductal papillomas, papillary cystadenomas are morphologically multicystic with numerous small to medium-sized cystic spaces. In the papillary cystadenoma the intraluminal growth is often characterized by multiple papillary projections with a variety of epithelial cell types, but usually the papillary growth occupies the lumen to a limited degree.

Prognosis and predictive factors
Excision appears to be curative based on five cases with an adequate follow-up of 2-5 years (1186,1302,1375,1829,2039).

Sialadenoma papilliferum

Definition
Sialadenoma papilliferum is an exophytic papillary and endophytic proliferation of mucosal surface and salivary duct epithelium.

ICD-O code 8406/0

Epidemiology
Sialadenoma papilliferum is a rare neoplasm (2711). The age range is 31-87 years (mean age 59) with a male to female ratio of 1.5:1 (264).

Localization
The vast majority of cases have occurred in the minor salivary glands. Major salivary gland involvement is very rare with the parotid gland being the most frequently involved. Over 80% of the neo-
Tumours of the salivary glands

Plasms occur on the hard and/or soft palate. Buccal mucosa is the second most common site. Other intraoral sites are the upper lip, the retromolar pad, and the faucial pillar.

Clinical features
The sialadenoma papilliferum typically manifests as a painless, exophytic papillary growth that is often interpreted clinically as a squamous papilloma. Duration ranges from months to several years.

Macroscopy
Gross findings usually show a well-demarcated papillary or verrucoid, sessile to pedunculated surface morphology. Overall, the tumours generally range from 0.5-1.5 cm in size.

Histopathology
The neoplasm consists of a biphasic pattern with a glandular component consisting of collections of cysts and duct-like spaces underlying a papillary or verrucous type proliferation of squamous epithelium. These papillary extensions of squamous epithelium are supported by fibrovascular cores and extend above the level of the adjacent mucosa. At or near the base of the fronds there is a transition from squamous epithelium to columnar ductal epithelium, which lines the proliferating ductal elements. These ductal elements consist of small and ectatic ducts, some of which show cystic enlargement. The ducts and their papillary folds are lined by a double row of cells showing a basal layer composed of cuboidal cells and a luminal lining of low columnar cells. Mucocytes can be interspersed throughout the lining of ductal cells as well as in the squamous component. Columnar oncotic cells may also be present.

Differential diagnosis
The differential diagnosis typically centres around three lesions: squamous papilloma, inverted ductal papilloma, and mucoepidermoid carcinoma. Squamous papilloma is composed entirely of squamous epithelium and lacks the endophytic growth pattern and glandular differentiation of sialadenoma papilliferum. Inverted ductal papilloma in contrast to the sialadenoma papilliferum, lacks the glandular complexity, and is a well-circumscribed tumour with blunted, pushing non-infiltrative margins. The invasive pattern and variable mixture of epidermoid, intermediate, mucous, and clear cells found in mucoepidermoid carcinoma set it apart from sialadenoma papilliferum.

Prognosis and predictive factors
The recurrence rate for sialadenoma papilliferum is in the 10-15% range based on 20 reported cases with adequate follow-up. Therefore, it is characterized by a higher risk of recurrence than the other types of ductal papillomas of the salivary gland. Complete surgical excision is the treatment of choice.
**Cystadenoma**

**Definition**
Cystadenoma is a rare benign epithelial tumour characterized by predominantly multicystic growth in which the epithelium demonstrates adenomatous proliferation. The epithelial lining is frequently papillary and rarely mucinous.

**ICD-O code**
8440/0

**Synonyms**
Monomorphic adenoma, cystic duct adenoma (2301), Warthin tumour without lymphoid stroma (668), intraductal papillary hyperplasia (401), oncocytic cystadenoma.

**Epidemiology**
The frequency of cystadenoma is between 4.2-4.7% of benign tumours (668,2711). There is a female predominance and the average age of patients with cystadenoma is about 57 years (range 12-89).

**Localization**
About 45% of all cases of cystadenoma arise in the parotid; the majority of tumours are located in minor salivary glands, particularly in the lips and buccal mucosa (668,2711).

**Clinical features**
Cystadenomas of the major glands typically present as slowly enlarging painless masses. In oral mucosa, these tumours produce smooth-surfaced nodules that resemble mucoceles.

**Macroscopy**
Cut section reveals multiple small cystic spaces or a single large cyst surrounded by lobules of salivary gland or by connective tissue.

**Histopathology**
Cystadenomas are often well circumscribed and surrounded by complete or incomplete fibrous capsules. The tumours are composed of cystic spaces, the number and size of which is variable. Twenty percent of cystadenomas are unilocular (2711). Most cases are multilocular with individual cystic spaces separated by limited amounts of intervening stroma. The lumens often contain eosinophilic material with scattered epithelial, inflammatory or foamy cells. Rarely, psammoma bodies or crystalloids have been described within the luminal secretion (2389). The lining epithelium of these cystic structures is mostly columnar and cuboidal. Oncocytic, mucous, epidermoid and apocrine cells are sometimes present focally or may even predominate. An oncocytic variant of cystadenoma is composed predominantly of oncocytes in unilayered or bilayered papillary structures thus resembling the epithelium of Warthin tumour without lymphoid stroma. Cystadenomas often show a mixture of cell types in the epithelial lining. An unusual case of oncocytic cystadenoma with apocrine, mucinous, sebaceous and signet ring cell appearance has been described (1715). Squamous epithelium may be present focally but rarely predominates. Cystadenomas of the salivary glands are usually devoid of foci of solid growth, cytologic atypia, fibrosis and apposed lymphoid tissue (790). Cystadenoma occurs in two major variants, as papillary and mucinous cystadenoma. Papillary cystadenoma is composed of large multilocular or unilocular cysts with multiple papillary projections. The lining epithelium is, in some cases, composed of oncocytic cells. Mucinous cystadenoma is composed of multiple cysts lined by mucous tall columnar epithelium with small basally situated nuclei and eosinophilic to clear cytoplasm. The lumens contain PAS and mucicarmine positive abundant mucus. The columnar epithelial lining has a uniform thickness with limited papillary growth.

**Prognosis and predictive factors**
Cystadenomas are benign tumours, and conservative but complete surgical removal is recommended. The tumours are unlikely to recur but rare cases of mucinous cystadenoma with malignant transformation have been described (1716).

**Fig. 5.74** Cystadenoma, composed of cystic spaces, the number and size of which is variable.
Fig. 5.75 Cystadenoma. A The lumen contains eosinophilic material. Cystic spaces are lined by columnar epithelium with focal apocrine metaplasia. B Scattered foamy cells within the secretion. C Oncocytic variant of cystadenoma is composed of prevailing oncocyes present in unilayered or bilayered papillary structures thus resembling Warthin tumour without lymphoid stroma. D Oncocytic cystadenoma with apocrine, mucinous, sebaceous and signet ring cell appearance.

Fig. 5.76 Cystadenoma. A Cystic spaces are lined by columnar epithelium with multiple papillary projections. B Prominent intracystic papillary growth pattern.
Excluding haematopoietic neoplasms, pure mesenchymal tumours account for 1.9-4.7% of salivary gland tumours [347, 669,678,2301] with benign soft tissue lesions being more common than sarcomas. The ratio of benign to malignant mesenchymal tumours varies from series to series, ranging from 18:1-2.4:1 (669,2301). Over 85% of soft tissue tumours arise in the parotid gland, over 10% involve the submandibular gland and, rarely, a tumour arises in the sublingual gland.

Vascular tumours are the most common benign mesenchymal neoplasm, accounting for almost 40% of the benign tumours [669,2301]. Seventy-five to 80% of the vascular neoplasms are haemangiomas, typically the juvenile or cellular variant, with the greatest incidence occurring in the first decade of life [430]. Most other vascular tumours are lymphangiomas. Other major salivary gland benign soft tissue neoplasms include neural tumours, most frequently neurofibroma or schwannoma [669] and fibroblastic/myofibroblastic tumours, most frequently nodular fasciitis, and fibromatoses with an infrequent myofibromatosis, fibroma, haemangiopericytoma, solitary fibrous tumour [953,2248] or inflammatory pseudotumour (inflammatory myofibroblastic tumour [775]) [2794] being reported. Lipomas, including the pleomorphic variety [934] and miscellaneous other tumours including granular cell tumour [2222], angiomyoma, glomangioma, myxoma, fibrous histiocytoma, giant cell tumour, osteochondroma and rarely a metastatic sarcoma may also be seen. Several cases of lipomas entrapping salivary glandular tissue have been recently described and termed sialolipomas [810,1824], including a congenital case [1129].

Salivary gland sarcomas arise in an older population than their benign soft tissue counterparts. They are rare tumours, accounting for only 0.3% of salivary gland neoplasms [347]. Almost any type of sarcoma can arise primarily in the salivary gland [87,669,1583]. In the largest published series, haemangiopericytoma, malignant schwannoma, fibrosarcoma and malignant fibrous histiocytoma were the most common neoplasms, accounting for 16.15, 14, and 11% of reported sarcomas, respectively [87]. These are aggressive neoplasms: 40-64% of patients develop recurrences, 39-64% develop metastases (usually haematogenous), and the mortality rate ranges from 36-64% with death occurring frequently within 3 years of diagnosis [87,1583]. The most successful treatments are wide surgical excision or surgery combined with radiation. For more specific information about each type of tumour, refer to the other texts [775,2745].

Table 5.04 Salivary gland sarcomas*

<table>
<thead>
<tr>
<th>Tumour type</th>
<th>Armed Forces Institute of Pathology Registry (87)</th>
<th>MD Anderson Medical Center (1583)</th>
<th>University of Hamburg Registry (2301)</th>
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<td>Malignant haemangioendothelioma</td>
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<tr>
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*Excluding lymphomas
**Arose in intraparotid lymph nodes
Haemangioma

**Definition**
This is characterized by a proliferation of endothelial cells and pericytes.

**ICD-O code**
9120/0

**Synonyms**
Benign or infantile haemangioendothelioma, infantile haemangioma, cellular haemangioma, immature capillary haemangioma, juvenile haemangioma

**Epidemiology**
Haemangiomas of the salivary glands account for approximately 0.4% of salivary tumours [668]. Lesions may present at any age but two thirds of cases are diagnosed in the first two decades [668,2301]. They are twice as common in females as males.

**Localization**
The haemangioma occurs almost exclusively in the parotid gland.

**Clinical features**
Lesions are asymptomatic soft swellings. They usually appear during the first 6 months of life and grow slowly. Most eventually involute by the age of 5-6 years [914,1413]. A bluish colour may be visible through skin but the overlying skin is not usually involved. Lesions are usually limited to the parotid gland but some are part of an angiomatosis that extends to involve the parapharyngeal space, infratemporal fossa or base of skull [1625]. Diagnosis may be aided by imaging [312,624].

**Macroscopy**
Lesions cause diffuse enlargement of the gland.

**Histopathology**
The lesion is composed of varying sized and shaped vascular spaces. The juvenile variant comprises small round densely packed endothelial cells and pericytes clustered within sheets that extend diffusely through the gland but divided into lobules by the gland septa. Lesional cells replace acinar cells, enlarging the lobules but leaving ducts scattered through the lesion. Mitoses are sparse or moderate in the juvenile form. In the early stages no vascular lumens are present but these develop with time to be the dominant feature [668,2301,2745]. Mature lesions are typical capillary haemangiomas with thin endothelial cell linings and no atypia. Thrombi and phleboliths may be present and foci of normal salivary tissue may persist in the mature lesion [668,914,1814,1817,2301].

**Prognosis and predictive factors**
Neonatal and infantile lesions grow rapidly initially but the majority involute before age 7 years and often much earlier [668,874,1413,1625,2301,2626,2632,2745,2790]. No treatment may be required and any intervention should be delayed. Steroids reduce growth and are the main treatment; pressure therapy [2626] or embolisation [265] may be considered if large. Occasional cases show progressive growth [2185].

![Fig. 5.78 Juvenile haemangioma. A Immature appearance with little lumen formation from a patient under 1 year in age. B More mature area with well-organised vessels. C Highly vascularized haemangioma.](image)
Hodgkin lymphoma

Involvement of the salivary glands by Hodgkin lymphoma is very rare. Combining data from four large series on primary lymphomas of the salivary glands, Hodgkin lymphoma only accounts for 4% of all cases \cite{473,893,1164,2267}. Both classical Hodgkin lymphoma and nodular lymphocyte predominant Hodgkin lymphoma have been reported \cite{101,391,893}, and all have involved the parotid gland only. Some of these tumours have originated from intraparotid lymph nodes, and thus are strictly-speaking not genuine primary extranodal lymphomas of the salivary glands. Rarely, Hodgkin lymphoma can arise within a Warthin tumour \cite{1702}. Please refer to the WHO Classification of Tumours of Haematopoietic and Lymphoid Tissues \cite{1197}.

Non-Hodgkin lymphoma

Overview

Primary salivary gland non-Hodgkin lymphomas (NHL) are uncommon, accounting for only 5% of all primary extranodal NHL \cite{809} and 2% of all salivary gland tumours \cite{893}. For a case to be considered as primary in the salivary gland, the bulk of disease should occur in this site, and the glandular parenchyma should be involved. A major problem in definition is caused by the normal presence of intraglandular lymph nodes in the parotid gland. Strictly speaking, cases of NHL limited to these lymph nodes without glandular parenchymal involvement should be considered as nodal NHL instead. However, the distinction is not always easy because cases with extensive parenchymal and nodal involvement can still have originated from intraglandular lymph nodes.

The most commonly affected gland is the parotid gland (75% of all cases), followed by the submandibular gland (20%) \cite{473,893,1164}. Most patients are in the sixth decade, and multiple glands (especially bilateral) are involved in about 10% of cases \cite{473}. The patients present with a palpable mass, and pain and tenderness are observed in a minority of cases.

Histologic types of NHL affecting the salivary glands

Most NHL occurring in salivary glands are B-cell lymphomas. In some older series, follicular lymphoma is the most common, accounting for about half of all cases \cite{473,893,1164}. However, many of these tumours are probably nodal lymphomas arising from intraglandular lymph nodes with subsequent infiltration of the glandular parenchyma, or represent extranodal marginal zone B-cell lymphoma of MALT type with prominent follicular colonization. In follicular lymphoma, lymphoepithelial lesions may be present in occasional cases \cite{1017}. Extranodal marginal zone B-cell lymphoma of MALT type is probably the most common type of lymphoma truly of salivary gland origin. It is frequently associated with Sjögren syndrome. Mantle cell lymphoma can also present as salivary gland involvement, but staging often reveals additional sites of disease. It is important not to misdiagnose mantle cell lymphoma for extranodal marginal zone B-cell lymphoma, because of the worse prognosis of the former.

Diffuse large B-cell lymphoma accounts for about 15% of all NHL of the salivary glands \cite{473,893,1164}. The tumour is infiltrative, with destruction of the salivary gland parenchyma and interstitial infiltration among residual salivary acini. The tumour comprises large lymphoid cells that may resemble centroblasts or immunoblasts, and express pan-B markers. Some cases represent transforma-
tion from an underlying extranodal marginal zone B-cell lymphoma of MALT type (2103). Anaplastic large cell lymphoma (ALCL), peripheral T-cell lymphoma unspecified, and extranodal NK/T cell lymphoma of nasal-type can also rarely affect the salivary glands (373,1081,1203). Please refer to the section of ‘non-Hodgkin lymphoma’ in ‘Tumours of the nasal cavity and paranasal sinuses’ for details.

Rare cases of NHL can arise in the lymphoid stroma of Warthin tumour, with follicular lymphoma being the most frequent type (307,1694). The lymphoma discovered in the Warthin tumour may be the presenting feature of more generalized disease.

Differential diagnosis

Some benign conditions can mimic NHL histologically. Lymphoepithelial sialadenitis (LESA), a condition associated with Sjögren syndrome, is a precursor lesion for extranodal marginal zone B-cell lymphoma of MALT type, and will be discussed in details in the next section. Kimura disease is a benign lesion of unknown etiology, commonly affecting the soft tissues in the head and neck region of young adults. It shows a predilection for Asian populations. Involvement of the major salivary glands is frequent (369,1385,1497,2581). It is characterized by reactive lymphoid follicles, vascularization of germinal centres, heavy eosinophilic infiltration, proliferation of high endothelial venules and prominent sclerosis. See Chapter 7 for details.

Chronic sclerosing sialadenitis (Küttner tumour) is a chronic inflammatory disorder affecting the submandibular gland (366). It can be bilateral. Since the gland is enlarged and hard, it usually imparts a clinical suspicion for malignancy. Histologically, there is a heavy lymphoplasmacytic infiltrate, accompanied by reactive lymphoid follicles, atrophy of salivary acini, periductal fibrosis and interlobular sclerosis. In the early phases, the striking lymphoid infiltrate can lead to a misdiagnosis of lymphoma. In contrast to lymphoma, the follicles are obviously reactive, the interfollicular lymphoid cells lack atypia, a permeative infiltrative pattern is lacking, there is a mixture of B and T cells, and the B cells are polytypic.

Rosai-Dorfman disease (sinus histiocytosis with massive lymphadenopathy) can also affect the major salivary glands (791,2760).

Prognosis and predictive factors

The prognosis of salivary gland lymphomas depends on the histologic type and clinical stage. T cell lymphomas and extranodal NK/T cell lymphomas are generally associated with a poorer outcome. A study reports that cases of probable nodal origin have a worse prognosis compared to those of probable extranodal-parenchymal origin (1193).

Extranodal marginal zone B-cell lymphoma (EMZBCL)

Definition

A low-grade B-cell lymphoma arising in mucosa-associated lymphoid tissue (MALT).

ICD-O code 9699/3

Epidemiology

Primary EMZBCL of the salivary gland is an uncommon neoplasm that usually develops in the setting of lymphoepithelial sialadenitis (LESA) in patients with Sjögren syndrome (59,98,710,2210,2266,2915). It occasionally occurs in the absence of an autoimmune disease or in association with another autoimmune process. EMZBCL occurs primarily in adults with an age range of 55-65 years and shows a slight female predominance. It may occasionally be seen in children and young adults (2524).

Etiology

Since most cases of EMZBCL arise in the
setting of LESA in association with Sjögren syndrome, it is postulated that this low-grade B cell lymphoma develops subsequent to the accumulation of mucosa-associated lymphoid tissue (MALT) that is acquired as a result of the autoimmune process.

**Localization**
EMZBCL usually presents as a persistent unilateral or bilateral mass in the parotid gland region, although any major or minor salivary gland may be involved (1181). The regional lymph nodes may also be enlarged due to involvement by the tumour.

**Clinical features**
EMZBCL may occur in the salivary gland as a manifestation of either primary or disseminated disease (473,813). Presenting signs include persistent enlargement of the involved salivary gland(s), sometimes in association with regional lymphadenopathy or monoclonal gammopathy (56). The patient may also show signs of other autoimmune diseases such as rheumatoid arthritis, systemic lupus erythematosus or Hashimoto thyroiditis (1181). Occasionally, EMZBCL may present in a cervical lymph node with subsequent development in the salivary gland. There is a variable period of time between the documented occurrence of LESA and the development of malignant lymphoma, which has been reported to range from 6 months to 29 years (630,710,2266). When EMZBCL of the parotid gland develops, there is a tendency for it to remain localized for long periods of time, as is the case with EMZBCL at other anatomic sites, including the stomach (468).

**Macroscopy**
The cut surface of EMZBCL of salivary gland is yellowish-tan in colour and has a “fish-flesh” appearance. Microcysts may be present.

**Tumour spread and staging**
The majority of patients with EMZBCL of salivary gland present with Stage IE (extranodal) or IIE disease. Dissemination most often occurs to cervical lymph nodes and other mucosal sites such as lung, conjunctiva and stomach (710,1003).

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Fig. 5.81 Extramedullary marginal-zone B-cell lymphoma (EMZBCL). A Lymphoepithelial lesion (left) highlighted by antibody to cytokeratin (right). B EMZBCL. Immunoglobulin light chain restriction for kappa (left) compared to lambda (right). C Follicular lymphoma of salivary gland with monotonous, neoplastic follicles extending into periglandular fat (left) and demonstrating reactivity for bcl-2 (right).
Histopathology
EMZBCL of the parotid gland occurs in a background of lymphoepithelial salivary adenitis (LESA) in almost all cases. The histologic features include a vaguely nodular to diffuse heterogeneous B-cell infiltrate that totally or subtotally effaces the normal glandular architecture. It is variably comprised of atypical small lymphocytes, centrocyte-like (cleaved) cells, monocytoid B-cells, immunoblasts, lymphoplasmacytic cells and plasma cells. Plasma cell differentiation may also be striking, causing confusion with a plasmacytoma. Intranuclear inclusions (Dutcher bodies) may be seen in the plasma or lymphoplasmacytic cells. Reactive germinal centres, often colonized by neoplastic B cells, are often present [1182]. Lymphoepithelial lesions, representing infiltration of the ductal and epithelial structures by neoplastic B cells, are seen in both LESA and EMZBCL. Ductal dilatation occasionally imparts a multicystic appearance to the gland. An important early change that occurs in EMZBCL of parotid gland, developing in the setting of LESA, is the formation of "halos", comprised of monocytoid and centrocyte-like B cells surrounding epithelial islands (lymphoepithelial lesion) [1162]. These cells may coalesce into broad, interconnecting sheets. Clusters of epithelioid histiocytes and prominent fibrosis may also be noted. There may be single or multifocal foci of large cell transformation adjacent to the low-grade component.

Immunoprofile
The B cell immunophenotype is confirmed by immunoreactivity for CD20 or CD79a. The lymphocytes and monocytoid B cells express surface immunoglobulin. The neoplastic B cells are negative for CD5, CD10, CD23 and bcl-1 (Cyclin D1). Bcl-2 reactivity in the neoplastic, colonizing B cells (but not in the residual, reactive germinal centre cells) is also characteristic. An antibody to cytokeratin may be useful to highlight the epithelial remnants in the lymphoepithelial lesions.

Differential diagnosis
The distinction between EMZBCL and LESA may be extremely difficult. Although histologic evaluation remains the gold standard for diagnosis, immunohistochemical, flow cytometric or molecular genetic analyses may be required. In both reactive follicular hyperplasia and EMZBCL, benign germinal centres are present but in the latter entity, the follicles may be colonized by neoplastic B-cells that express bcl-2. A dense diffuse B cell infiltrate, intranuclear inclusions (Dutcher bodies) and cytologic atypia are characteristically seen in EMZBCL. Lymphoepithelial lesions may be seen in both LESA and EMZBCL. The demonstration of light chain restriction by immunohistochemistry or flow cytometry supports the monoclonality of the B cell lymphoma. Extramedullary marginal zone lymphoma with prominent nodularity may simulate a follicular lymphoma (FL). It is necessary, therefore, to distinguish the reactive, colonized germinal centres in EMZBCL from the neoplastic germinal centres in FL. The majority of cases of FL will show immunoreactivity for bcl-2 and will express the germinal centre cell markers CD10 and bcl-6.

Somatic genetics
There are clonal rearrangements of the immunoglobulin genes {2,104,608,630, 2103,2524}. The significance of this finding, however, is somewhat unclear and controversial since gene rearrangements have also been found in histologically benign cases of LESA [105,770,2103] and in the salivary gland lesions of Sjögren syndrome patients who subsequently developed overt lymphoma [1238]. The cell of origin of EMZBCL lymphoma has not been definitively identified and has been postulated to be of post germinal centre origin. In some cases, however, the variable regions of the immunoglobulin genes have been shown to undergo somatic hypermutation, suggesting that this tumour may arise from germinatal centre B cells [104]. Although no specific oncogene has been described in association with MALT lymphoma, numerical chromosomal abnormalities, especially Trisomy 3 [293,2823, 2824], and the chromosomal translocation, t (11;18) (q21;q21) {1961} have been reported in EMZBCL in various anatomic sites.

Prognosis and predictive factors
The prognosis of salivary gland EMZBCL lymphoma is usually very favourable. Tumours that are localized (Stage IE) at the time of presentation and demonstrate purely low-grade histology have an excellent prognosis. With lymph node involvement (Stage IIE), the prognosis is usually similar to primary nodal low-grade B cell lymphomas. Although EMZBCL of salivary gland may show histologic transformation to a higher grade, similar to what has been reported in the stomach, the clinical significance of this finding remains unclear. There is no conclusive evidence that treatment of EMZBCL prevents transformation to a higher grade. With or without treatment, EMZBCL of the salivary gland is usually an indolent neoplasm that does not result in significant morbidity or mortality. There are reports, however, of patients with EMZBCL associated with LESA subsequently developing extensive extra salivary gland lymphoma or nodal large B-cell lymphoma.

Salivary gland extramedullary plasmacytoma
Please refer to Chapter 1 for details.
Secondary tumours

Definition
A metastatic tumour involving salivary glands that originates in a distant site.

Epidemiology
Secondary tumours comprise about 5% of all malignant tumours of salivary glands (669,2293,2300), but this incidence is considerably higher in some countries (199). The peak incidence is in the 7-8th decade. Almost 70% of cases occur in males. The majority of cases are squamous cell carcinoma and melanoma is second in frequency.

Localization
The large majority of metastases are located in the parotid, while fewer are seen in the submandibular gland. Metastases occur within the interstitial tissue and the intra-/periglandular lymph nodes with a slight predominance of extranodal infiltrates (2300).

Clinical features
Eighty percent of secondary tumours of the parotid are metastases from head and neck neoplasms. On the other hand, 85% of metastatic tumours in the submandibular glands are from distant sites (899). Primary sites frequently are the upper and middle parts of the facial region (including skin, mucous membranes, deep soft tissues as well as eyes and ears) (443,692,1917,1987). A further 10% originate from distant tumours among which lung carcinoma (especially the small cell carcinoma type), kidney and breast carcinomas are the most common (669,806,1585,2219,2293, 2300). However, almost 10% of the secondary tumours remain undefined as to their origin.

Histopathology
Generally, metastases retain to some extent the histological pattern and cytological characteristics of the respective primary tumour.

Fig. 5.82 Small cell carcinoma diffusely infiltrating the salivary gland tissue. Tumour cells are marked by immunoreactivity to synaptophysin (right).

Fig. 5.83 Metastatic renal cell carcinoma in the parotid gland. Low magnification.
Odontogenic tumours and tumour-like lesions constitute a group of heterogeneous diseases that range from hamartomatous or non-neoplastic tissue proliferations to benign neoplasms to malignant tumours with metastatic potential. They are derived from epithelial, ectomesenchymal and/or mesenchymal elements of the tooth-forming apparatus. Odontogenic tumours are rare, some even extremely rare, but can pose a significant diagnostic and therapeutic challenge.
### MALIGNANT TUMOURS

**Odontogenic carcinomas**
- Metastasizing (malignant) ameloblastoma\(^1\)  
  9310/3
- Ameloblastic carcinoma – primary type  
  9270/3
- Ameloblastic carcinoma – secondary type (dedifferentiated), intraosseous  
  9270/3
- Ameloblastic carcinoma – secondary type (dedifferentiated), peripheral  
  9270/3
- Primary intraosseous squamous cell carcinoma – solid type  
  9270/3
- Primary intraosseous squamous cell carcinoma derived from keratocystic odontogenic tumour  
  9270/3
- Primary intraosseous squamous cell carcinoma derived from odontogenic cysts  
  9270/3
- Clear cell odontogenic carcinoma  
  9341/3
- Ghost cell odontogenic carcinoma  
  9302/3

**Odontogenic sarcomas**
- Ameloblastic fibrosarcoma  
  9330/3
- Ameloblastic fibrodentin– and fibro-odontosarcoma  
  9290/3

### BENIGN TUMOURS

**Odontogenic epithelium with mature, fibrous stroma without odontogenic ectomesenchyme**
- Ameloblastoma, solid / multicystic type  
  9310/0
- Ameloblastoma, extraosseous / peripheral type  
  9310/0
- Ameloblastoma, desmoplastic type  
  9310/0
- Ameloblastoma, unicystic type  
  9310/0
- Squamous odontogenic tumour  
  9312/0
- Calcifying epithelial odontogenic tumour  
  9340/0
- Adenomatoid odontogenic tumour  
  9300/0
- Keratocystic odontogenic tumour  
  9270/0

**Odontogenic epithelium with odontogenic ectomesenchyme, with or without hard tissue formation**
- Ameloblastic fibroma  
  9330/0
- Ameloblastic fibroodontoma  
  9290/0
- Ameloblastic fibrodentinoma  
  9271/0
- Odontoma, complex type  
  9282/0
- Odontoma, compound type  
  9281/0
- Odontoameloblastoma  
  9311/0
- Calcifying cystic odontogenic tumour  
  9301/0
- Dentinogenic ghost cell tumour  
  9302/0

**Mesenchyme and/or odontogenic ectomesenchyme with or without odontogenic epithelium**
- Odontogenic fibroma  
  9321/0
- Odontogenic myxoma / myxofibroma  
  9320/0
- Cementoblastoma  
  9273/0

**Bone-related lesions**
- Ossifying fibroma  
  9262/0
- Fibrous dysplasia  
- Osseous dysplasias  
- Central giant cell lesion (granuloma)  
- Cherubism  
- Aneurysmal bone cyst  
- Simple bone cyst

**OTHER TUMOURS**
- Melanotic neuroectodermal tumour of infancy  
  9363/0

\(^1\) Morphology code of the International Classification of Diseases for Oncology (ICD-O) (821) and the Systematized Nomenclature of Medicine (http://snomed.org).

Behaviour is coded /0 for benign tumours, /3 for malignant tumours, and /1 for borderline or uncertain behaviour.
Odontogenic tumours are lesions derived from epithelial, ectomesenchymal and/or mesenchymal elements that still are, or have been, part of the tooth-forming apparatus. These tumours, therefore, are found exclusively within the maxillofacial skeleton (intraosseous or centrally located), or in the soft tissue (gingiva) overlying tooth-bearing areas or alveolar mucosa in edentulous regions (extraosseous or peripherally located). The tumours may be generated at any stage in the life of an individual. Knowledge of basic clinical features such as age, gender, and location can be extremely valuable in developing differential diagnoses of odontogenic tumours.

Prior consensus conferences on taxonomy of odontogenic tumours, cysts and allied lesions [2048,2050,2051,2148,2595} confirmed that the characteristic morphological and inductive relationship between the various parts of the normal tooth germ are reproduced, to a greater or lesser extent, in many of the tumours and tumour-like lesions of the odontogenic tissues. The observation of these features is important both in the identification of the lesions and in their classification. For example, normal dentin is easily identified because of its tubular structure, but if for some reason this tubular structure is absent it is difficult to distinguish between atypical poorly mineralized dentin (dentinoid) and atypical osteoid. However, if an osteoid-like tissue develops in direct juxtaposition to odontogenic epithelium, this relationship provides presumptive evidence that the material is dysplastic dentin.

The classification used here is based firstly on a lesion’s behaviour, with a classification into benign, malignant and non-neoplastic. Subdivisions of “benign” lesions are then based on the types of odontogenic tissues involved: odontogenic epithelium with mature, fibrous stroma without odontogenic ectomesenchyme; odontogenic epithelium with odontogenic ectomesenchyme, with or without hard tissue formation; mesenchyme and/or odontogenic ectomesenchyme with or without the presence of odontogenic epithelium.

**Epidemiology**

Data from China, Hong Kong, Nigeria, Zimbabwe, Germany, Turkey, Japan, Canada, South Africa and the US show marked differences in relative frequencies between benign odontogenic tumours. According to the PRC study, the most frequent tumour was ameloblastoma, solid/multicystic type (A-S/M, 58.6%) comparable to that found in Hong Kong (59.4%), Japan (57%) and in two African countries (Zimbabwe and Nigeria), 79.1% and 58.5%, respectively. This contrasts with the rates in series involving populations in the US and Canada, where the most frequent tumour was odontoma (73.8% and 56.4%, respectively) with A-S/M accounting only for 12.2% and 14.8%, respectively. It seems that one reason for these discrepancies may be found in the source of the data. Odontogenic tumour patients from PRC, Hong Kong, Japan and several African countries are diagnosed and treated in Maxillofacial Units of Medical Hospitals, whereas patients from the US and Canada generally are monitored in Dental Schools or Hospitals. Odontomas frequently are diagnosed on routine panoramic images performed in a dentist’s surgery or in a Dental School without previous biopsy. In several developing countries cases are not registered or sent for histological confirmation. Thus, the frequency of odontomas reported from these countries is probably underestimated. Ameloblastomas (A-S/Ms) on the other hand need a biopsy for confirmation of the diagnosis, and the radical treatment is often performed in a Medical Hospital. As it has been suggested that ameloblastomas are more common in Blacks than in Caucasians, it remains to be proved that the geographical variation suggested by the above data may also be based on ethnic differences. Benign odontogenic neoplasms (including hamartomatous lesions) seem to outnumber their malignant counterparts by a factor as high as 100 (531).

**Etiology**

The etiology of benign and malignant odontogenic tumours is unknown. The majority of odontogenic tumours seem to arise de novo, without an apparent causative factor.

**Clinical features**

The large majority of odontogenic tumours occurs intraosseously within the maxillofacial skeleton, while extraosseous odontogenic tumours occur nearly always in the tooth-bearing mucosa. The clinical features of most benign odontogenic tumours are non-specific; benign odontogenic tumours show slow expansive growth with no or slight pain. In contrast, pain is the first and most common symptom followed by rapidly developing swelling in nearly all malignant odontogenic tumours. The tumour may erode or break through the cortex of the jaw bones.
odontogenic tumours cross-sectional imaging studies (CT, MRI) cover both topography and fine structure of the lesion, and give valuable information about tumour extension.

**Precursor lesions**
Developmental odontogenic cysts may contribute to the formation of certain odontogenic tumours and intraosseous squamous cell carcinoma.

**Odontogenesis and genetics**
Odontogenesis depends on the sequential and reciprocal interactions between cranial neural crest (CNC)-derived ectomesenchymal cells and the epithelium that lines the oral cavity (357). Contact of these CNC-cells with oral epithelium initiates tooth development by the formation of an epithelial tooth bud with a surrounding condensation of CNC cells. It is not yet clear whether the initial odontogenic potential lies in CNC cells inducing the epithelium to form the tooth bud or the converse, CNC cells responding to an inductive stimulus from the oral epithelial lining (2897).

Experimental and family cluster studies of the molecular events associated with tooth development have resulted in the identification of over 200 genes playing a role in this area (464). Of these, the fibroblast growth factor–8 partitions the jaw into an alveolar and a basal part, whereas sonic hedgehog is involved with the formation of the tooth bud (464). Expression of both these genes in turn induces upregulation of additional genes, both in epithelium and ectomesenchyme as odontogenesis proceeds.

**Classification**
Malignant odontogenic tumours (MOTs) are classified as odontogenic carcinomas and odontogenic sarcomas. Odontogenic carcinomas include in the previous WHO classification (2051) and more recently discussed (2394) is not included in the present classification due to lack of evidence for its existence as an entity.

**Epidemiology**
Generally, MOTs are rare entities. Some of them are exceedingly rare. Odontogenic carcinomas seem to occur more frequently in the elderly (2402).

**Etiology**
The etiology of MOTs is unknown.

**Clinical features / Imaging**
Clinical symptoms are identical to those of other malignant tumours in the maxillofacial region. Swelling, pain, bleeding, ulceration of the oral mucosa, mobility of teeth, paraesthesia or anaesthesia may be indicators of a MOT. Involvement of local lymph nodes and distant metastases may occur early in the course of the disease. Extensive jaw bone destruction with ill defined borders are characteristic and in addition a mixed radiographic pattern may be evident. Hard tissue structures may be diagnosed on plain radiographs, CTs or MRIs.

**Therapy**
Due to the small number of published cases of MOTs specific treatment protocols or guidelines are at present not available. Surgical resection with tumour-free margins is the therapy of choice.

**Precursor lesions**
In some cases MOTs have been demonstrated to develop from preexisting benign counterparts.

**Prognosis and predictive factors**
Generally, the prognosis for MOTs is poor.

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**Fig. 6.2** Dental anlage with initial formation of dental hard structures (<). EM, ectomesenchyme; OM, oral mucosa. RDL, rests of dental lamina; SR, stellate reticulum; OEE, outer enamel epithelium; IEE, inner enamel epithelium.

**Malignant odontogenic tumours**

**Definition**
Most malignant odontogenic tumours are generally considered the counterparts of benign odontogenic tumours, but others such as the primary intraosseous squamous cell carcinoma are not.
Metastasizing ameloblastoma

Definition
Metastasizing ameloblastoma is an ameloblastoma that metastasizes in spite of a benign histologic appearance.

ICD-O code
Metastasizing (malignant) ameloblastoma 9310/3

General features
Metastasizing ameloblastoma shows no specific features different from ameloblastomas that do not metastasize. Therefore, this diagnosis can only be made in retrospect, after the occurrence of metastatic deposits. Thus, it is clinical behaviour and not histology that justifies a diagnosis of metastasizing ameloblastoma. Ameloblastomas with atypia are ameloblastic carcinomas (1875). Confusion may also arise through the use of the term atypical ameloblastoma to denote lesions with fatal outcome for various reasons, either metastasis, histological atypia or relentless local spread (50). Metastatic deposits of ameloblastomas are mostly seen in the lung (1386,1439, 2405) but have also been reported at other sites.

Ameloblastic carcinoma – primary type

Definition
The ameloblastic carcinoma is a rare primary odontogenic malignancy that combines the histological features of ameloblastoma with cytological atypia. This will be the case even in the absence of metastases.

ICD-O code 9270/3

Epidemiology
The incidence of ameloblastic carcinoma is unknown, fewer than 60 cases have been reported. The reason for the high number of reported cases from China where 6.7% of all odontogenic tumours are malignant is not clear (1569).

Clinical features / Imaging
Approximately 2/3 of ameloblastic carcinomas involve the mandible (492,1813). Only 19 cases have been reported to occur in the maxilla (593). Males and females are equally affected. The posterior segments of the jaws represent the most common site. Generally, ill defined or irregularly margined radiolucencies are characteristic. Cortical expansion often with perforation, may be present as well as infiltration into adjacent structures

Histopathology
Ameloblastic carcinoma is characterized by malignant cytologic features in combination with the overall histological pattern of an ameloblastoma. A tall columnar cellular morphology with pleomorphism, mitotic activity, focal necrosis, perineural invasion and nuclear hyperchromatism may be present. Peripheral palisading and so-called reverse or inverted nuclear polarity will be present. A stellate reticulum structure will usually be seen. Cystic spaces may be present that are lined by epithelium (497). Atypical cells form nests and broad ribbons which may branch and anastomose with focal areas of subtle necrosis to more obvious central, comedo necrosis-like areas. Ameloblastic carcinomas show a high proliferation index compared to benign ameloblastomas by
virtue of an increased index of proliferating cell nuclear antigen in addition to higher levels of aneuploidy (1315,1793).

Differential diagnosis
The differential diagnosis includes ameloblastomas which may show an occasional mitotic figure. Other odontogenic carcinomas including primary intraosseous squamous cell carcinoma and clear cell odontogenic carcinoma are usually distinct; however, some ameloblastic carcinomas have been reported to show clear cell features and yet others may contain a spindle cell component. Metastatic carcinomas should also be considered; however, they do not display ameloblastic features.

Somatic genetics
Aneuploidy has been found to be more frequently present in the ameloblastic carcinoma and may be used as a strong predictor of malignant potential of questionble lesions (1793). More recently 5q13 amplification was demonstrated by comparative genomic hybridization (CGH) in an ameloblastic carcinoma (1289).

Prognosis and predictive factors
Maxillary ameloblastic carcinomas demonstrate tumour-related deaths or pulmonary metastases in over one-third of cases (593). Mandibular counterparts behave in a similar manner, where local recurrences are likely to precede metastases (2363).

**Ameloblastic carcinoma – secondary type (dedifferentiated), intraosseous**

**Definition**
Ameloblastic carcinoma arises in a pre-existing benign ameloblastoma. The term “dedifferentiated ameloblastoma” has been applied when morphologic features of typical ameloblastoma were noted (1029,2405,2651). This in turn separates metastazising ameloblastoma since cytologic atypia is not a feature of this entity.

**ICD-O code** 9270/3

**Synonym**
Carcinoma ex intraosseous ameloblastoma

**Epidemiology**
Dedifferentiated ameloblastoma or ameloblastic carcinoma arising within a pre-existing microscopically verified benign ameloblastoma is a very rare occurrence. Most cases arise in older individuals (7th decade) (1029,2651), usually with a clinically proven long-standing ameloblastoma.

**Ameloblastic carcinoma – secondary type (dedifferentiated), peripheral**

**Definition**
Transformation of a pre-existing peripheral extraosseous ameloblastoma to a malignant cellular phenotype (1673,1674). Prior cases of so-called intraoral basal cell carcinomas (gingiva) may, in retrospect, be considered within this category as well (102,635,2033,2780).

**ICD-O code** 9270/3

**Synonym**
Carcinoma ex peripheral ameloblastoma

**Epidemiology**
To date six cases of an ameloblastic car-
cinoma arising within a preexisting peripheral ameloblastoma with a 1:1 gender distribution have been reported [2033].

**Clinical features/Imaging**
A gingival mass with variable surface alterations including irregularity, concavity, sessile, and pedunculated features as well as alveolar bone resorption have characterized the previously reported cases of transformed peripheral ameloblastoma. They are generally non-tender, and may be associated with a developing inter-radicular radiolucency with separation of the roots of adjacent teeth [102,1674].

**Histopathology**
Nests, strands and follicles of recognizable ameloblastoma-type histology within the gingival soft tissues may be noted in association with variable degrees of squamous differentiation. An extensive network of tumour islands with peripherally located columnar cells and stellate reticulum-type areas will be present. Nests of keratin, cellular and nuclear pleomorphism, and abnormal mitotic figures will extend to and invade alveolar bone and around peripheral nerves [320].

**Prognosis and predictive factors**
Wide local excision with en bloc resection of the involved segment of the affected jaw bone is the treatment of choice. Long-term follow-up is mandatory [2780].

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Fig. 6.6 Ameloblastic carcinoma, secondary type. Aggregates of moderately differentiated squamous cell carcinoma abutting ameloblastoma with peripheral palisading of tumour cells and central parts reminiscent of stellate reticulum.
Primary intraosseous squamous cell carcinomas

Primary intraosseous squamous cell carcinoma - solid type

Definition
Primary intraosseous squamous cell carcinoma (PIOSCC) is a central jaw carcinoma derived from odontogenic epithelial remnants. Subcategories of PIOSCC include (1) a solid tumour that invades marrow spaces and induces osseous resorption, (2) squamous cancer arising from the lining of an odontogenic cyst and (3) a squamous cell carcinoma in association with other benign epithelial odontogenic tumours. When the tumour destroys the cortex and merges with the surface mucosa, it may be difficult to distinguish between a PIOSCC and a true carcinoma arising from the oral mucosa. Invasion from an antral primary must also be excluded.

ICD-O code 9270/3

Synonym
Primary intra-alveolar epidermoid carcinoma

Epidemiology
The male: female ratio approaches 2:1, with a mean age of 55 years, although cases have been encountered during infancy.

Etiology
These tumours arise centrally in the jaws, with no communication with the upper aerodigestive tract mucosa and are therefore not subjected to exposure of the usual carcinogenic factors.

Localization
PIOSCC is more often found in the body and posterior mandible than the maxilla. Maxillary cases are most frequently seen in the anterior segment.

Clinical features
Signs and symptoms
Most cases are asymptomatic and are discovered incidentally during the course of routine dental radiographs. Facial swelling may be observed. Perineural invasion of the inferior alveolar nerve will produce paresthesia of the lip.

Imaging
Radiographically, PIOSCCs are osteolytic. The margins of the radiolucency are often irregular and noncorticated. Larger extensive lesions may show cortical bone expansion and destruction.

Tumour spread and staging
PIOSCC spreads both regionally and distantly.

Macroscopy
Gross features of PIOSCCs are those of any carcinoma within bone.

Histopathology
PIOSCC is characterized by islands of neoplastic squamous epithelium with the features of squamous cell carcinoma. Most lesions are moderately differentiated without prominent keratinization. The stroma may or may not exhibit an inflammatory infiltrate. Metastatic squamous cell carcinoma has to be excluded. There are no specific histopathologic features that distinguish a metastatic squamous cancer from PIOSCC. When the central jaw carcinoma can be documented histologically to arise from the epithelial lining of an odontogenic cyst, primary site of origin is accepted. Mucoepidermoid carcinoma also arises from the lining of odontogenic cysts and must be included in the differential diagnosis. Special staining for mucin may be helpful.

Histogenesis
Carcinomas that arise centrally within the jaw bones putatively arise from epithelial remnants of odontogenesis that include the periradicular rests of Malassez and
the reduced enamel epithelium surrounding impacted teeth [2397]. Rarely, a central jaw squamous cell carcinoma represents dedifferentiation from a benign ameloblastoma [2651].

**Primary intraosseous squamous cell carcinoma derived from keratoctytic odontogenic tumour**

**Definition**
A squamous cell carcinoma arising within the jaws without connection to the oral mucosa in the presence of an keratoctytic odontogenic tumour (KCOT).

**ICD-O code**
9270/3

**Epidemiology**
Thirteen cases of primary intraosseous squamous cell carcinoma deriving from KCOT have been documented [1305, 1615]. Most of these lesions are encountered in older patients, 40 years and above, with male predilection.

**Etiology**
There are no known specific predisposing factors.

**Localization**
The mandible is involved much more frequently than the maxilla [1305].

**Clinical features / Imaging**
Early lesions are generally insidious [1305,1615], usually presenting as a benign odontogenic cyst and the diagnosis of carcinoma is only made after microscopic examination. In others, local symptoms relate to the effects of the lesion at specific sites i.e. pain, swelling, loosening of teeth, non-healing extraction sockets, and paraesthesia. Lesions in advanced disease stage frequently appear as overtly malignant growths with associated ulceration and induration. Regional lymphadenopathy may also be present.

Radiographically, early lesions are often indistinguishable from any odontogenic cyst. In some, the margins of the radiolucent defect appear irregular and ‘ragged.’ Late lesions are obviously destructive. A multilocular appearance with cortical destruction and frequent soft tissue extension characterizes these late-stage neoplasms.

**Histopathology**
The histological appearance of this lesion is typically that of a keratinizing well-differentiated squamous cell carcinoma in conjunction with KCOT. The main differential diagnosis would include keratoameloblastoma, squamous odontogenic tumour, central high-grade mucoid carcinoma and metastatic lesions [1305].

**Genetics**
One case of mandibular odontogenic carcinoma revealed abnormalities in a small subset of genes [34].

**Prognosis and predictive factors**
Lack of information precludes definitive prognostication.

cases. In a series of 28 patients the mean age was 56 years and the male to female ratio was almost 2:1.

**Clinical features / Imaging**
The majority occur in the mandible with symptoms of pain, paraesthesia or anaesthesia of the lower lip. The radiographic aspect may mimic any type of odontogenic cyst.

**Histopathology**
Histopathologically, the tumour is characterized as a cyst lined by any type of epithelium that can be seen in odontogenic cysts in association with a squamous cell carcinoma. Various degrees of dysplasia may be observed in the epithelial cyst lining. The architecture of verrucous hyperplasia or verrucous carcinoma may be present, as well.

**Prognosis and predictive factors**
PIOSCC associated with an impacted lower third molar seems to have a favourable prognosis; however, the number of reported cases is small.

**Primary intraosseous squamous cell carcinoma derived from odontogenic cysts**

**Definition**
A squamous cell carcinoma arising within the jaws without connection to the oral mucosa, and in the presence of an odontogenic cyst other than keratoctytic odontogenic tumour.

**ICD-O code**
9270/3

**Epidemiology**
There are less than fifty well-documented
Clear cell odontogenic carcinoma

Definition
Clear cell odontogenic carcinoma (CCOC) is characterized by sheets and islands of vacuolated and clear cells.

ICD-O code 9341/3

Historical annotation
In the past CCOC was called clear cell ameloblastoma [2712] and clear cell odontogenic tumour [699,1004] and was considered a benign tumour in the previous WHO classification of 1992 [2051].

Epidemiology
Only 36 cases have been reported during the last 15 years. The tumour has a strong female predilection and tends to occur in older adults, the mean age at diagnosis being close to 60 years, range 17-89 [1784].

Localization
The most frequently affected site is the mandible.

Clinical features / Imaging
The CCOC usually causes swelling of the jaws and loosening of teeth. Aggressive tumour growth results in an ill-defined radiolucency, and root resorption may occur [114].

Histopathology
A biphasic pattern is often exhibited. CCOC is primarily composed of a fibrous stroma with islands of epithelial cells revealing clear to faintly eosinophilic cytoplasm, well-demarcated cell membranes and irregular dark-staining nuclei. Also cords of dark-staining basaloid cells with scant eosinophilic cytoplasm may be seen [114]. In addition, ameloblastomatous islands with palisaded peripheral cells may be observed. Mitoses and necrosis are rare. Histochemically, many of the tumour cells contain abundant delicate and coarse diastase degradable PAS-positive granules, but they are negative for mucin and amyloid [114,521].

Immunoprofile
The clear and eosinophilic tumour cells are consistently reactive for cytokeratins 13, 14, 19, 8, 18 and EMA. They are negative for vimentin, S-100-protein, desmin, smooth muscle actin, HMB-45, alpha (1)-chymotrypsin, CD31, CD45 and GFAP [114,521,699,1172,1499].

Differential diagnosis
Since clear cells frequently may be seen in other neoplasms in the oral and maxillofacial region, it is important to rule out lesions like salivary gland tumours, melanotic tumours, metastatic renal cell carcinoma and clear cell variant of calcifying epithelial odontogenic tumour [1172,1784,2040].

Genetics
DNA-analysis has shown a polyploid population with DNA index of 1.93 and an S-phase of 10.2% [114]. Comparative genomic hybridization discloses consistent chromosomal aberrations in both primary and metastatic CCOC [275].

Prognosis and predictive factors
The CCOC exhibits an aggressive growth pattern and frequently recurs. The tumour can metastasize to regional lymph nodes and lungs, as well as to bone, and tumour progression may even cause tumour related death. Consequently, resection with tumour-free margins is the treatment of choice, and long-term follow-up is mandatory. Adjuvant radiotherapy is a rational option for tumours that have eroded cortical bone [256].

Fig. 6.10 Clear cell odontogenic carcinoma. A Biphasic tumour pattern with sheets of clear cells and irregular cords and strands of dark, basaloid cells, intersected by narrow bands of fibrous stroma. B Sheets of polygonal cells with central clear or faintly eosinophilic cytoplasm, well defined cell membrane and dark, peripheral or central nucleus. Peripheral cells abutting on narrow fibrous bands are palisaded. Tumour cells show clear cytoplasm with apical nucleus.
Ghost cell odontogenic carcinoma

Definition
Ghost cell odontogenic carcinoma is a malignant odontogenic epithelial tumour with features of calcifying cystic odontogenic tumour and/or dentinogenic ghost cell tumour.

ICD-O code 9302/3

Synonyms
Calcifying ghost cell odontogenic carcinoma, malignant epithelial odontogenic ghost cell tumour, carcinoma arising in a calcifying odontogenic cyst, aggressive epithelial ghost cell odontogenic tumour, malignant calcifying odontogenic cyst and malignant calcifying ghost cell odontogenic tumour.

Epidemiology
Only 19 cases have been reported in the English literature and more than half of them were from Asia. The tumour occurs more commonly in men than women (male: female=2:1) and affects patients in the age range of 13-72 years with a peak incidence in the fourth decade.

Localization
The tumour occurs more commonly in the maxilla than the mandible (2:1), either at anterior or posterior area, corresponding to the site distribution of the calcifying odontogenic cyst.\(^\{297}\).

Clinical features / Imaging
Clinical features include swelling, often with paraesthesia. Imaging shows a poorly demarcated, osteolytic radiolucency with some radiopaque material. Displacement of tooth roots is common and tooth impaction and root resorption are occasionally noted. Large lesions in the maxilla often destroy the sinus wall, grow into the nasal and orbital cavities, and extend to adjacent structures.

Macroscopy
A typical lesion consists of a well-circumscribed cystic portion and a solid portion with gritty consistency on cut-surface, although some are completely solid.

Histopathology
The diagnosis of the neoplasm is based on the identification of a malignant epithelial tumour containing classic benign features of calcifying cystic odontogenic tumour. The malignant component consists of rounded epithelial islands in a fibrous stroma. The epithelial cells are either small, rounded with dark nuclei or larger with vesicular nuclei. Many mitoses are seen. Ghost cells are found in varying numbers either isolated or in clusters. Dysplastic dentin may be present\(^{1314,1568}\). The relationship of the benign and malignant features appears to have two distinct forms. In the first pattern, the malignant epithelial component is physically separated from the classical benign lesion, which is either cystic or solid. The other pattern is an admixture of the malignant epithelial component with typical benign features\(^{672,1260,1568}\). PCNA labelling index is a possible parameter in differentiating the ghost cell odontogenic carcinoma from its benign counterparts. PCNA labelling indices (65.9±7.3% and 65.2±5.6%) in the malignant epithelial cells are significantly higher than in the benign neoplastic (45.8%) and cystic variants (29.3% and 11.6%)\(^{1460,2536}\). Immunohistochemical overexpression of p53 protein is demonstrated in the tumour cells\(^{1568}\).

Prognosis and predictive factors
The prognosis is unpredictable due to a wide spectrum of growth patterns. They vary from a slowly growing, locally invasive tumour to a highly aggressive and rapidly growing neoplasm with local recurrence and metastasis. The overall five-year survival rate is 73%. Recurrences are common\(^{341,1260,1314,1568,2536}\).
**Odontogenic sarcomas**

**Ameloblastic fibrosarcoma**

**Definition**
Ameloblastic fibrosarcoma (AFS) is an odontogenic tumour with a benign epithelial and a malignant ectomesenchymal component. It is regarded as the malignant counterpart of the ameloblastic fibroma (AF).

**ICD-O code**
9330/3

**Synonym**
Ameloblastic sarcoma

**Epidemiology**
There is a wide age range (3-89 years) with a mean age of 27.5 years at diagnosis, versus the AF (14.8 years) [2034]. Patients with AFS derived from a preexisting AF have a mean age of 33 years [1794]. Those with de novo AFS have a mean age of 22.9 years [266]. Sixty three percent of reported cases have occurred in males and 37% in females.

**Etiology**
The etiology of AFS is unknown. Approximately one third of AFS represents malignant transformation of a preexisting AF.

**Localization**
The mandible is the most commonly affected site (78%), followed by the maxilla (20%) [266]. In both jaws the posterior region is the site of predilection. Only one case of peripheral AFS has been published.

**Clinical features / Imaging**
Typically, AFS presents as an expansile intraosseous radiolucency with ill-defined borders [2394]. Swelling and pain are common findings. Paraesthesia has been observed [266].

**Macroscopy**
AFS has a fleshy consistency, with a white to yellowish cut surface.

**Histopathology**
The histologic pattern of AFS resembles ameloblastic fibroma in which the epithelial tissue is benign but the connective tissue component is malignant. The epithelium is composed of budding and branching cords of small polygonal epithelial cells admixed with islands and knots. Larger islands have a border of columnar cells with hyperchromatic nuclei. A hypercellular connective tissue stroma displaying mitotically active cells surrounds the epithelial component [2394]. Recurrent tumours tend to show greater stromal cellularity and mitotic rate. In addition, the amount of the epithelial component may decrease or disappear [2394, 2539].

**Prognosis and predictive factors**
The biologic behaviour of AFS is that of a highly locally aggressive neoplasm with extremely low potential for distant metastasis. Of 64 cases only one had metastasis to mediastinal nodes and liver [266, 440, 532, 917, 1794].

**Ameloblastic fibrodentino – and fibro-odontosarcoma**

**Definition**
A tumour with histological features of ameloblastic fibrosarcoma, together with dysplastic dentin (fibro-dentinosarcoma) and/or enamel/enameloid and dentin/dentinoid (fibro-odontosarcoma).

**ICD-O code**
9290/3

**Synonyms**
Ameloblastic dentinosarcoma; ameloblastic odontosarcoma; ameloblastic sarcoma; odontogenic sarcoma.

**Epidemiology**
To date only fourteen cases have been reported in the literature [266, 1794, 2539]. Nine cases occurred in men and four in women. The age range is 12-83 years with a peak in the third decade.

**Clinical features / Imaging**
Most cases present a slow growing pain-
less jaw swelling. Radiographically, the lesions are radiolucent, sometimes multilocular with poorly circumscribed outlines. One or more dense opacities representing the hard tissue components may be present.

Precursor lesions
Ameloblastic fibro-odontomas are regarded as possible precursor lesions [46,1135,1794].

Histopathology
The tumour shows the typical features of ameloblastic fibrosarcoma in addition to the formation of dental hard tissues in scattered areas [46,1462,2540,2814].

Prognosis and predictive factors
The treatment of choice is surgical resection [46,2394]. Ameloblastic sarcomas seem to have a better prognosis than other jaw sarcomas and can generally be regarded as low-grade. Only one case has shown regional metastases. Local recurrences are more often seen [2540].

Fig. 6.14 Ameloblastic fibrosarcoma. A At low magnification, the tumour has an ameloblastic fibroma-like pattern. B Nests and cords of odontogenic epithelium in a highly cellular odontogenic ectomesenchyme. C Marked pleomorphism and mitotic figures in odontogenic ectomesenchyme adjacent to an epithelial island.
Ameloblastomas

Ameloblastoma, solid / multicystic type

Definition
The solid/multicystic ameloblastoma (A-S/M) is a slowly growing, locally invasive, epithelial odontogenic tumour of the jaws with a high rate of recurrence if not removed adequately, but with virtually no tendency to metastasize.

ICD-O code 9310/0

Synonyms
Conventional ameloblastoma; classical intraosseous ameloblastoma.

Epidemiology
Although rare, the A-S/M is the second most common odontogenic tumour. It exhibits no gender predilection and occurs over a wide age range. Most cases are diagnosed between 30 and 60 years of age, while the tumour is rare below the age of 20 years. Geographic and racial differences have been described [1431,2325].

Etiology
The cause of A-S/M is not known. Dysregulation of several genes in normal tooth development may play a role in its histogenesis [1048].

Localization
The tumour occurs exclusively in the jaws, rarely in the sinonasal cavities. Approximately 80% occur in the mandible, with a marked predilection for the posterior region, except in African Blacks in whom any region of the mandible may be involved, particularly the symphysis [427]. Most maxillary examples occur in the posterior region.

Clinical features / Imaging
Small A-S/Ms may be asymptomatic. More commonly, A-S/Ms present as variably sized swellings of the jaws. Pain or paraesthesia are rare. A-S/Ms may be unilocular or multilocular radiolucencies resembling cysts and they may reveal scalloped borders. An unerupted tooth may be associated with A-S/M. Resorption of the roots of adjacent teeth is common. Definitive diagnosis of A-S/Ms cannot be made radiologically, since similar radiographic features are displayed by e.g. keratocystic odontogenic tumour or myxoma. Particularly for maxillary A-S/Ms, CT-scans or MRIs are recommended.

Tumour spread and staging
A-S/Ms spread slowly by infiltration through the medullary spaces and may erode cortical bone. Eventually, it will resorb the cortical plate and may extend into adjacent tissues. Tumours of the posterior maxilla tend to obliterate the maxillary sinus and subsequently extend intracranially.

Histopathology
There are two basic histopathologic patterns, the follicular and plexiform, without clinical relevance. The follicular pattern consists of islands of odontogenic epithelium within a fibrous stroma. Typically, the basal cells of these islands are columnar, hyperchromatic, and lined up in a palisaded fashion. Typically their nuclei are displaced away from the basement membrane, and their cytoplasm is generally vacuolated. The central cells may be loosely arranged, resembling...
stellate reticulum. These areas often become cystic and at times confluent. If these cells are spindle-shaped, basaloid, granular or showing squamous differentiation, the terms spindle cell ameloblastoma, basal cell ameloblastoma, granular ameloblastoma and acanthomatous ameloblastoma have been used. In addition ghost cells may be observed. The plexiform pattern contains basal cells arranged in anastomosing strands with an inconspicuous stellate reticulum. The stroma is usually delicate, often with cyst-like degeneration. For both histologic patterns, mitotic activity and cellular pleomorphism are rarely noted. The microscopic differential diagnosis may include ameloblastic fibroma, squamous odontogenic tumour, adenomatoid odontogenic tumour, odontogenic remnants in dental follicles, epithelial-rich odontogenic fibroma, calcifying odontogenic cyst and adenoid cystic carcinoma arising form the maxillary sinus.

Genetics
A recent study using comparative genomic hybridization described chromosomal aberrations in 2 of 17 A-S/Ms (1191). A notably homogeneous gene profile in eight examples has been shown by cDNA array. Thirty-four of the 588 genes studied, demonstrated differences in ameloblastomas compared to tooth germs. The fos-oncogene and tumour-necrosis-factor-receptor-1 (TNFRSF-1A) were the most over-expressed genes. Ten genes, including sonic hedgehog (SHH), cadherins 12 and 13 (CDH12, and 13), plus transforming growth-factor-ß1 (TGF-ß1), were underexpressed in all ameloblastomas studied (1048).

Prognosis and predictive factors
Long-term follow up is essential, since recurrences have been noted more than ten years after the initial treatment. Treatment should include excision with an adequate margin of uninvolved tissues. Lesions involving the posterior maxilla, demonstrate the poorest prognosis. Radiotherapy should not be used in the first line treatment of A-S/M. Metastasizing ameloblastoma is discussed elsewhere in this volume (page 287).

Ameloblastoma, extraosseous / peripheral type

Definition
The extraosseous / peripheral amelo-
blastoma (A-E/P) is the extraosseous counterpart of the intraosseous solid / multicystic ameloblastoma (A-S/M).

**ICD-O code** 9310/0

**Synonyms** Soft tissue ameloblastoma, ameloblastoma of mucosal origin, ameloblastoma of the gingiva.

**Epidemiology** Extraosseous/peripheral ameloblastomas (A-E/Ps) comprise 1.3-10% of all ameloblastomas [2033]. Age range varies from 9 and 92 years with 64% of all cases occurring in the fifth through seventh decade. The mean age of patients with A-E/Ps (males: 53 years; females: 51 years) is significantly higher than for the intraosseous counterpart which has a mean age of 37 years [2149]. The male:female ratio is 1.9:1 [2149].

**Localization** A-E/Ps is located to the tooth-bearing areas (gingiva) or alveolar mucosa in edentulous areas. A mandible:maxilla ratio of 2.4:1 is noted. Multicentric origin of A-E/Ps has been reported [1075].

**Clinical features / Imaging** The A-E/P is a painless, firm and exophytic growth with a smooth, pebbly or papillary surface. Rarely, intraosseous ameloblastomas may extend to the gingival tissues and merge with the gingival epithelium, creating an exophytic A-E/P-like lesion [2473]. Apart from a superficial erosion or depression (saucerization or cupping) of the bone crest due to pressure resorption, there is rarely significant bone involvement [2149].

**Macroscopy** The gross specimen consists of a firm to spongy, pinkish-grey tissue mass.

**Histopathology** The A-E/P consists of odontogenic epithelium with the same histomorphological cell types and patterns as seen in A-S/M. Some lesions are located entirely within the connective tissue of the gingiva, showing no continuity with the surface epithelium, whereas others seem to fuse with or originate from the mucosal epithelium. It is generally believed that basal cell carcinoma of the gingiva (BCCG) and A-E/P represent the same neoplasm [861]. Squamous cells in the acanthomatous areas of A-E/Ps may show ghost cell formation, and in some parts of the tumour islands, vacuolated or clear cells occur in discrete clusters [1879,2137]. The stroma is that of a mature, fibrous connective tissue. Rare cases of malignant A-E/Ps (ameloblastic carcinomas) have been reported [2033, 2526,2649].

**Differential diagnosis** Differential diagnosis includes: (1) peripheral odontogenic fibroma. The proliferation of strands and islands of odontogenic epithelium in this tumour may be so extensive as to make the distinction from A-E/P difficult [862], (2) peripheral variant of the squamous odontogenic tumour [2029], and (3) odontogenic gingival epithelial hamartoma (OGEH) [103]. The present view is that A-E/P and OGEH represent the same lesion [2033].

**Histogenesis** A-E/P may arise from odontogenic epithelial remnants within the gingival lamina propria or from the basal cell layer of the gingival epithelium.

**Prognosis and predictive factors** A-E/P does not show invasive behaviour and conservative excision is the treatment of choice. The recurrence rate is low (16-19%) [300,1852]. Long-term follow-up is recommended.
**Epidemiology**
A-D is similar to A-S/M regarding age and gender distribution [2035,2352].

**Localization**
The maxilla: mandible ratio is 1:1. The ratio for A-S/M is 1:5.4 [2149]. The A-Ds are found predominantly in the anterior mandibular region.

**Clinical features / Imaging**
A painless swelling of the jaw bone represents the chief initial complaint. The size of the tumour varies between 1.0 and 8.5 cm in diameter. An extraosseous variant of A-D has not been reported. Radiographically, about 50% of A-Ds show a mottled, mixed radiolucency/radiopacity with diffuse margins, suggesting a fibro-osseous lesion. Resorption of tooth roots and bone formation may occur. The ill-defined borders of A-Ds make high-resolution CT and MRI helpful in treatment planning [2595].

**Macroscopy**
The lesional tissue has a gritty consistency; the cut surface is solid in most cases.

**Histopathology**
In A-Ds the stromal component dominates, compressing the odontogenic epithelial components. The epithelial tumour islands are very irregular or bizarre in shape with a pointed, stellate appearance. The epithelial cells at the periphery of the islands are cuboidal with occasional hyperchromatic nuclei. Columnar cells with nuclear polarity are rarely conspicuous. The islands have a swirled, hypercellular centre with spindle-shaped or squamous, epithelial cells. Microcysts may occur centrally. Myxoid changes of the juxtaepithelial stroma are often found. Formation of metaplastic osteoid trabeculae (osteoplasia) may be present [2035]. A fibrous capsule is not present corresponding to the radiographically poorly defined tumour margin. A combination of A-D with A-S/M is known and has been termed as “hybrid lesion” [1703,2035].

**Immunoprofile**
In contrast to A-S/M, marked immunostaining of TGF-β has been observed [2537].

**Prognosis and predictive factors**
Present knowledge leads to the recommendation to apply the same treatment modality as for A-S/M.

**Ameloblastoma, unicystic type**

**Definition**
The unicystic ameloblastoma (A-U) represents an ameloblastoma variant, presenting as a cyst.

**ICD-O code**
9310/0

**Synonym**
Cystogenic ameloblastoma

**Epidemiology**
Cases associated with an unerupted tooth show a mean age of 16 years as opposed to 35 years in the absence of an unerupted tooth [2030]. The mean age is significantly lower than that for A-S/M. There is no gender predilection [13]. Five to 15% of all ameloblastomas are of the unicystic type [2149].

**Localization**
More than 90% of cases involve the mandible, usually the posterior region [13].

**Clinical features / Imaging**
Some cases are asymptomatic, sometimes presenting as a swelling of the posterior mandible. Up to 80% are associated with an unerupted mandibular third molar. The lesion presents radiographically as a well corticated unilocular, often pericoronal radiolucency [702,1461]. Root resorption may occur [2149]. The clinical radiographic diagnosis is frequently a dentigerous (follicular) cyst.

**Fig. 6.21** Unicystic ameloblastoma. Panoramic radiograph mimicking dentigerous (follicular) cyst with impacted second mandibular molar.

**Fig. 6.22** Schematic view of histological variants of unicystic ameloblastoma: luminal (ameloblastomatous cyst epithelium), intraluminal (protruding into cyst cavity) and mural (left and right, invading cyst wall).

**Fig. 6.23** Intraluminal variant of unicystic ameloblastoma with plexiform pattern.
Macrosopy
The lesions vary in size and, when removed intact, are typically cystic, and generally attached to an unerupted tooth at the cemento-enamel junction. The cyst wall may contain one or more tumour proliferations extending into the lumen. These proliferations and other thickened areas must be selected for microscopic examination.

Tumour spread and staging
The A-U is an expansile lesion that can destroy a significant portion of the jaw. The A-U does not usually behave as an A-S/M and does not infiltrate the surrounding bone.

Histopathology
Two histopathologic variants exist. The luminal variant is a cystic lesion lined by ameloblastomatous epithelium. In addition intraluminal extensions may occur. These extensions usually exhibit a plexiform epithelial pattern. There is no tumour infiltration into the fibrous wall. The mural variant, the cyst wall is infiltrated by ameloblastomatous epithelium that exhibits either a follicular or plexiform pattern. Sometimes both variants may occur in the same lesion. The mural variant of A-U may be confused with either dentigerous cysts or dental follicles containing a lot of odontogenic epithelial remnants. These epithelial nests, however, do not show the typical histologic features of ameloblastoma; peripheral palisading and nuclear polarization.

Prognosis and predictive factors
Most A-Us are enucleated with the preoperative clinical diagnosis of dentigerous cyst and it is only on pathologic examination that their true nature is determined.

WULIN 6.24 Unicystic ameloblastoma (luminal type) showing ameloblastomatous epithelium lining the cyst wall.
Squamous odontogenic tumour

Definition
Squamous odontogenic tumour (SOT) is a locally infiltrative neoplasm consisting of islands of well-differentiated squamous epithelium in a fibrous stroma.

ICD-O code
9312/0

Epidemiology
The SOT is a rare tumour with less than 50 cases reported. The age range is between 8-74 years with a mean of 38.7 years (1170). The gender ratio is 1.4:1 (men:women).

Etiology
The etiology of SOT is unknown.

Localization
SOT usually occurs intraosseously and probably develops in the periodontal ligament between the roots of vital erupted, permanent teeth. The mandible is affected more often than the maxilla.

Clinical features / Imaging
Mobility of teeth, local pain, swelling of the gingiva, osseous expansion or mild gingival erythema may be observed (979). Radiographically, a unilocular or triangular radiolucency between the roots of adjacent teeth is seen. Extensive SOTs may present a multilocular pattern. The peripheral variant may produce some ‘saucerization’ of the underlying bone – a result of pressure from tumour expansion rather than neoplastic infiltration.

Histopathology
SOTs are composed of islands of well-differentiated squamous epithelium of varying size and shape. Islands are rounded or oval, but irregular and cord-like structures may be seen. Individual tumour islands reveal a peripheral layer of low cuboidal or flat epithelial cells. Epithelial islands may undergo central microcystic degeneration. Individual islands may contain calcified material. SOTs may be mistaken for an acanthomatous ameloblastoma, a desmoplastic ameloblastoma, a well-differentiated squamous cell carcinoma, or pseudoepeitheliomatous hyperplasia. SOT must be differentiated from so-called “squamous odontogenic tumour-like islands arising in the walls of odontogenic cysts”.

Prognosis and predictive factors
Conservative surgical treatment is usually sufficient. Recurrences are rare and are probably due to incomplete removal.
Calcifying epithelial odontogenic tumour

Definition
The calcifying epithelial odontogenic tumour (CEOT) is a locally invasive epithelial odontogenic neoplasm, characterized by the presence of amyloid material that may become calcified.

ICD-O code
9340/0

Synonym
Pindborg tumour [2046,2047].

Epidemiology
CEOT accounts for approximately 1% of all odontogenic tumours occurring in patients between 20 and 60 years of age, with a mean around 40 years [2031]. There is no gender predilection. Most cases are intraosseous, approximately 6% arise in extraosseous locations. Intraosseous tumours affect the mandible more often than the maxilla with a ratio of 2:1. There is a predilection for the premolar/molar region, although any site may be involved. Peripheral lesions usually occur in the anterior gingiva [1133].

Clinical features / Imaging
The tumour presents as an asymptomatic slow-growing expansile mass of the jaw. Peripheral gingival lesions are firm painless masses. Radiographically, most CEOTs present as mixed radiolucent-radiopaque lesions, but they may show considerable variation. They may be unilocular or multilocular. In about half of the cases, an unerupted tooth, most often a mandibular third molar, is associated with the lesion. Computed tomography and magnetic resonance imaging provide useful information in the diagnosis and treatment of CEOT [507].

Macroscopy
Macroscopic features are those of a solid tumour with various amounts of calcification. Cystic change is not seen.

Histopathology
The tumour consists of a fibrous stroma with islands and sheets of polyhedral epithelial cells with abundant eosinophilic cytoplasm, sharply defined cell borders and well-developed intercellular bridges. Their nuclei are frequently pleomorphic, with giant nuclei being common. Mitotic figures are rarely encountered; in case of malignant transformation, mitoses are frequent [2688].

Eosinophilic, homogeneous hyalin material that is often calcified in the form of concentric rings is present within or around the sheets of tumour cells. Positive staining with Congo red and fluorescence with thioflavine T show this material to be amyloid. Some tumours are amyloid-rich, while others demonstrate epithelial-predominance. Calcification is characteristic but non-calcifying variants also occur [2399,2538]. Lack of calcification is more common in extraosseous tumours. Clear cells may be present within the epithelial nests; they contain glycogen. In some cases, the clear cells make up a significant proportion of the tumour [2031]. In a minority of cases, the CEOT may be seen in a composite relationship with the adenomatoid odontogenic tumour [2031].

Differential diagnosis
Due to the presence of cytonuclear pleomorphism and intercellular bridges, CEOT may be mistaken for intraosseous...
squamous cell carcinoma, either primary or metastatic. The clear cell variant of the CEOT must be distinguished from clear cell odontogenic carcinoma, metastatic carcinomas composed of clear cells such as renal cell carcinoma and salivary gland malignancies including mucoepidermoid carcinoma and acinic cell carcinoma.

**Prognosis and predictive factors**
The CEOT is a locally invasive tumour. Small tumours may be enucleated, but larger ones require local resection. An overall recurrence rate of about 14% has been noted (803). A relatively higher recurrence rate of 22% has been noted for the clear cell variant (1086). Long-term follow-up is recommended.

Fig. 6.31 CEOT. Giant and pleomorphic nuclei in the absence of mitoses are frequently found and do not indicate malignancy. Note intracellular homogenously dispersed eosinophilic material representing amyloid.

Fig. 6.32 Calcifying epithelial odontogenic tumour. Congo red staining shows green birefringence when subjected to polarized light.

Fig. 6.33 Calcifying epithelial odontogenic tumour. Histochemical findings of the hyalin-like material. The material is positively stained with Congo red.
Adenomatoid odontogenic tumour

Definition
Adenomatoid odontogenic tumour (AOT) is composed of odontogenic epithelium in a variety of histoarchitectural patterns, embedded in a mature connective tissue stroma and characterized by slow but progressive growth.

ICD-O code 9300/0

Epidemiology
The AOT accounts for 2-7% of all odontogenic tumours (2028, 2032). The age range varies between 3 and 82 years. More than two thirds are diagnosed in the second decade of life and 90% are found before the age of 30. More than half of the cases occur among teenagers (2032). The male:female ratio is 1:1.9. In some Asian countries the ratio may reach 1:3.2 (2613).

Localization
The AOT almost exclusively occurs intraosseously with a preference for the maxilla over the mandible with a ratio of 2.1:1. The rare peripheral type occurs almost exclusively in the anterior maxillary gingiva (2036).

Clinical features / imaging
Intraosseous AOTs may be found in association with unerupted permanent teeth (follicular type), in particular the four canines that account for 60% with the maxillary canines alone accounting for 40%. Most AOTs are asymptomatic. When growth of the intraosseous variants causes cortical expansion, it may present as a palpable bony-hard swelling with or without slight pain. The intraosseous AOTs may cause displacement of neighbouring teeth. The peripheral variant presents as a fibroma or an epulis-like lesion of the gingiva.
Radiographically, the intraosseous, follicular AOT, shows a well-defined, unilocular radiolucency around the crown and often part of the root of an unerupted permanent tooth, mimicking a dentigerous cyst. If not associated with an unerupted tooth (extrafollicular type), AOT presents as a unilocular radiolucent lesion. In two thirds of the intraosseous variant, the radiolucency shows discrete radiopaque foci (2036). The peripheral variant may disclose erosion (saucerization) of the alveolar bone crest.

Histopathology
At low magnification the most striking pattern is that of variably sized solid nodules of cuboidal or columnar cells of odontogenic epithelium forming nests or rosette-like structures with minimal stromal connective tissue. Between the epithelial cells and in the centre of the rosette-like configurations, eosinophilic amorphous material (“tumour droplets”) is present. Conspicuous within the cellular areas are structures of tubular or duct-like appearance. The duct-like spaces are lined by a single row of columnar epithelial cells, with the nuclei polarized away from the luminal surface. The duct-like spaces represent pseudolumina formed by secretion of the columnar epithelial cells. The lumen may be empty or contain eosinophilic material or cellular debris. The duct-like structures may not be present in all AOTs. In addition to forming ducts, the cuboidal to columnar cells form convoluted cords in complicated patterns that often exhibit invaginations. Another characteristic cellular pattern is composed of nodules consisting of polyhedral, eosinophilic epithelial cells of squamous appearance with distinct cell boundaries and prominent intercellular bridges. The nuclei may occasionally reveal mild (degenerative) pleomorphism. These nodules may contain pools of amorphous amyloid-like material and globular masses of calcified substances. Melanin pigmentation of both lesional tissue and stroma cells has been described. Occurrence of a hyaline, dysplastic material or calcified osteodentin...
may be found in AOTs. It is likely the result of a metaplastic process, as odontogenic ectomesenchyme is not present, and thus should not be interpreted as an induction phenomenon, although in very rare cases dentin-like material containing dentinal tubules may occur. The mature connective tissue stroma of the AOT is generally loosely structured and contains thin-walled congested vessels. CEOT-like areas found in AOTs should be considered a histological variant of AOT, as should areas of AOTs mimicking calcifying ghost cell odontogenic cysts (2888), developing odontomas or other odontogenic tumours or hamartomas.

**Prognosis and predictive factors**

AOTs are cured by local excision. Recurrences are extremely rare [2032].

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**Fig. 6.35** Adenomatoid odontogenic tumour. AOT with a solid nodule of cuboidal epithelial cells (SN) containing several eosinophilic, amorphous “tumour droplets” (arrows), a duct-like structure (DS) lined by a single row of columnar epithelial cells. At the periphery strands of epithelium in a cribriform pattern.

**Fig. 6.36** Adenomatoid odontogenic tumour (AOT). Solid, cell-rich area of minimal stromal connective tissue showing duct-like structures (arrows), and convoluted structure (CS) of tall columnar epithelial cells.

**Fig. 6.37** Adenomatoid odontogenic tumour combined with calcified epithelial odontogenic tumour AOT/CEOT. The tumour shows areas of CEOT-like foci (asterisks) in an otherwise typical AOT with cribriform configurations of epithelial strands and several dilated thin-walled vessels in the sparse stroma.
Definition
A benign uni- or multicystic, intraosseous tumour of odontogenic origin, with a characteristic lining of parakeratinized stratified squamous epithelium and potential aggressive, infiltrative behaviour. It may be solitary or multiple. The latter is usually one of the stigmata of the inherited naevoid basal cell carcinoma syndrome (NBCCS).

ICD-O code 9270/0

Synonyms
The traditional designation is odontogenic keratocyst (OKC), which stresses the benign behaviour of this lesion. However, the WHO Working Group recommends the term keratocystic odontogenic tumour (KCOT) as it better reflects its neoplastic nature. Other synonyms include odontogenic keratocystoma and primordial cyst.

Epidemiology
Keratocystic odontogenic tumours (KCOT) occur from the first to the ninth decades with a peak in the second and third decades [2323]. The mean age of patients with multiple KCOTs, with or without the NBCCS, is lower than those with single non-recurrent KCOTs. Most series have shown a preponderance in males.

Etiology
Recent studies have demonstrated the role of the PTCH gene in the etiology of KCOTs [110,143,471,1344,1467,1468,1481,1854].

Localization
The mandible is involved more frequently than the maxilla, with figures ranging from 65-83% of cases. About one-half originate at the angle of the mandible, extending anteriorly and superiorly [2323].

Clinical features
The most important clinical feature of the KCOT is its potential for locally destructive behaviour, its recurrence rate, and its tendency to multiplicity, particularly when associated with the NBCCS. Patients may complain of pain, swelling or discharge. These tumours may reach a large size prior to discovery. KCOT may penetrate cortical bone and involve adjacent structures.

Imaging
KCOTs may appear as small, round or ovoid unilocular radiolucencies or may be larger with scalloped margins. A mandibular radiolucency may involve body, angle and ascending ramus. The radiolucencies tend to be well-demarcated with distinct sclerotic margins, but may be diffuse in parts. Maxillary lesions tend to be smaller, but more extensive involvement may occur. True multilocular mandibular lesions are not uncommon.
Adjacent teeth may be displaced but root resorption occurs rarely [2323].
CT scans may be helpful in detecting cortical perforation and assessment of soft tissue involvement. They may be of particular value in the evaluation of patients with multiple NBCCS-related KCOTs [1563]. Contrast enhanced MRI may provide more detailed information [1204, 1744].

**Macroscopy**
Linings are thin and fragile, and are usually collapsed and folded.

**Histopathology**
The KCOTs are lined by a regular parakeratinized stratified squamous epithelium, usually about 5-8 cell layers thick and without rete ridges. There is a well-defined, often palisaded, basal layer of columnar or cuboidal cells. The nuclei of the columnar basal cells tend to be oriented away from the basement membrane and are often intensely basophilic. This is an important feature in distinguishing KCOT from jaw cysts with keratinization. The parakeratotic layers often have a corrugated surface. Desquamated keratin is present in many of the cavities. Mitotic figures are found frequently in the suprabasal layers. Some linings may show features of epithelial dysplasia [22] but malignant transformation to squamous cell carcinoma is rare [1600].
In the presence of an intense inflammatory process, the epithelial lining loses its characteristic cellular and architectural features.
Cystic jaw lesions that are lined by orthokeratinizing epithelium do not form part of the spectrum of a keratocystic odontogenic tumour (KCOT).

**Histogenesis**
It is generally agreed that KCOT arises from odontogenic epithelium. The available evidence points to two main sources of the epithelium: the dental lamina or its remnants and extensions of basal cells from the overlying oral epithelium [284,2323,2478].

**Genetics**
The NBCCS or PTCH gene has been mapped to chromosome 9q22.3-q31 [720] and probably functions as a tumour suppressor. Studies on NBCCS and sporadic KCOT have provided molecular evidence of a two-hit mechanism in the pathogenesis of these tumours demonstrating allelic loss, at two or more loci, of 9q22 [1481,1539] leading to overexpression of bcl-1 and TP53 in the NBCCS. This supports the concept that KCOT represents a neoplasm [1539]. There is also accumulating evidence that the PTCH gene might be a significant factor in the development of sporadic KCOT [143,1344,1467,1468,1854]. Furthermore, preliminary results have shown over-expression and amplification of genes located in 12q [1047].

**Prognosis and predictive factors**
The KCOT is a potentially aggressive lesion. Patients should be carefully followed up after treatment because of the common presence of daughter cysts and a tendency for multiplicity.
**Ameloblastic fibroma / fibrodentinoma**

**Definition**
Ameloblastic fibroma (AF) consists of odontogenic ectomesenchyme resembling the dental papilla and epithelial strands and nests resembling dental lamina and enamel organ. No dental hard tissues are present. If there is dentin formation, the lesion is referred to as ameloblastic fibrodentinoma (AFD).

**ICD-O code**
- Ameloblastic fibroma: 9330/0
- Ameloblastic fibrodentinoma: 9271/0

**Epidemiology**
AF is a rare odontogenic tumour (531, 1930, 2140). The mean age is 14.8 years (range from 7 weeks to 62 years).

**Localization**
AF mainly occurs in the posterior mandible (2034).

**Clinical features / Imaging**
Most cases of AF present as painless swelling or are discovered due to disturbances of tooth eruption. Radiographically, the tumour presents as a well-demarcated radiolucency, often in connection with a malpositioned tooth (2034).

**Histopathology**
The epithelial component of AF consists of branching and anastomosing epithelial strands that form knots of varying size. These have a peripheral rim of columnar cells similar to the inner enamel epithelium that embraces a loosely arranged spindle-shaped epithelium identical to stellate reticulum. The epithelial strands lie in a myxoid cell-rich stroma with stellate-shaped fibroblasts with long slender cytoplasmic extensions resembling embryonic tooth pulp. The amount of epithelium may vary. Dental hard tissues do not form part of the histologic spectrum of AF. Mitotic figures both in epithelial and mesenchymal components may occur; if present, they should raise concern about the benign nature of the case.

The epithelial component resembles ameloblastoma. The stromal component however differs in that it is an immature cell-rich myxoid tissue with an embryonic appearance. Some AFs may contain granular cells (2034, 2539). Rarely, tumours with the histomorphology of AF may form dysplastic dentin, and are called ameloblastic fibrodentinomas (AFD) (2051, 2539). Histologic features similar to AF may be observed in the hyperplastic dental follicle (1313, 2489). The distinction relies on correlation of the histology to the radiographic appearance: a radiolucent rim surrounding an unerupted tooth, in case of a dental follicle and an expansive radiolucent jaw lesion, in case of ameloblastic fibroma.

**Prognosis and predictive factors**
Treatment consists of enucleation and curettage. Recurrence may occur but this does not justify initial aggressive treatment (2034). Rarely, AF may progress to malignancy (ameloblastic fibrosarcoma).
Ameloblastic fibro-odontoma

Definition
Ameloblastic fibro-odontoma (AFO) is a tumour, which has the histologic features of ameloblastic fibroma (AF) in conjunction with the presence of dentin and enamel.

ICD-O code 9290/0

Epidemiology
AFO is less common than AF. The mean age is between 8-12 years. There is no gender or anatomic site predilection.

Clinical features / Imaging
AFO is often asymptomatic and may be detected as a result of failure of tooth eruption. Radiographically, AFO exhibits a well-circumscribed unilocular or multilocular radiolucency with varying levels of radiopacity depending on the extent of mineralization. AFO is often associated with an unerupted tooth.

Histopathology
AFO is composed of soft and hard tissues. The soft tissue component is identical to AF; the hard tissue component consists of dental hard structures. These two components may be present in varying proportions. Odontoameloblastoma may be considered in the differential diagnosis.

Prognosis and predictive factors
The prognosis is excellent; recurrences have been rarely described.

Fig. 6.45 Ameloblastic fibro-odontoma. A Ameloblastic fibro-odontoma of the left mandible showing radiopacities in a large translucency. The second molar is displaced and impacted. B CT scan of the same patient.

Fig. 6.46 Ameloblastic fibro-odontoma. A Well-formed dentin and enamel in the lesion. B Initial induction of dentin- and enamel-matrices in the proliferating soft tissue similar to that of ameloblastic fibroma.
Odontoma, complex type

Definition
Odontoma, complex type (OC) is a tumour-like malformation (hamartoma) in which enamel and dentin, and sometimes cementum, is present.

ICD-O code
Odontoma 9280/0
Odontoma, complex type 9282/0

Synonym
Complex composite odontoma

Epidemiology
OC is one of the most common odontogenic tumours. It is primarily diagnosed in children, adolescents, and young adults \{1104,2034,1004\}. There is no gender predilection.

Etiology
The etiology is unknown \{2037\}.

Localization
OCs occur in tooth-bearing regions, mostly in the posterior part of the mandible.

Clinical features / Imaging
OCs are painless slowly growing lesions. Growth stops when they are fully matured some reaching up to 6 cm in diameter. The majority measures less than 3 cm. Swelling of the jaw may be evident. Adjacent teeth may be displaced, and impaction of a permanent tooth is a common finding \{1949A\}. Radiographically, OCs appear as a spherical or ovoid radiopacity with a fine radiating periphery, surrounded by a radiolucent zone, which may be broader in a developing complex odontoma. Differential diagnosis from a compound odontoma or even an osteoma may not be possible radiographically.

Histopathology
In mature OCs the soft tissue capsule consists of a loose connective tissue containing strands or islands of odontogenic epithelium. In developing OCs the outer part of the odontoma consists of a cell rich zone of soft tissue with formation of dentin and enamel, not resembling tooth morphology. The lesion appears as a mass of primarily tubular dentin which encloses hollow circular or oval structures with empty spaces from decalcified mature enamel, enamel-matrix producing epithelium and connective tissue. The structure of the hard dental tissue may vary. The lesion consists mainly of wavy and plicated walls of tubular or dysplastic dentin covered by enamel. Between these walls are irregular curvilinear clefts that contain enamel matrix-producing epithelium and connective tissue. Cementum is scarce except on the “root” surfaces of tooth-like structures. Scattered ghost cells may be present \{2288A\}. The distinction between complex and compound odontoma is mainly based on the presence of tooth-like structures in compound odontomas. The differential diagnosis between a developing complex odontoma and an ameloblastic fibro-odontoma is sometimes impossible.

Prognosis and predictive features
Complex odontomas are treated by local excision. Recurrences have only been reported in cases of incomplete removal of developing complex odontomas \{2167A\}.

Fig. 6.47 Odontoma, complex type, with a dense radiopacity and a retained molar.

Fig. 6.48 Odontoma, complex type. Enamel, dentin, and cementum-like tissue are arranged in a haphazard pattern, in contrast to the regular structure encountered in compound odontoma.
Odontoma, compound type

Definition
A tumour like malformation (hamartoma) with varying numbers of tooth-like elements (odontoids).

ICD-O code
Odontoma 9280/0
Odontoma, compound type 9281/0

Synonym
Compound composite odontoma

Epidemiology
Odontoma, compound type (OCp) is primarily diagnosed in children and adolescents [1104,2034] with no gender predilection. It has been reported to be the most common of all odontogenic neoplasms and tumour-like lesions.

Etiology
The etiology is unknown [2037].

Localization
OCps may occur in any tooth-bearing area of the jaws. The anterior maxilla is most frequently affected.

Clinical features / Imaging
OCps are painless, slowly growing lesions. When fully matured, development ceases. The size usually varies between 1 and 2 cm in diameter, but a diameter of up to 6 cm has been reported. Swelling of the jaw is seen in less than 10% of the cases. Many are located close to the incisal/occlusal part of an impacted tooth, thus impeding eruption. Displacement of erupted teeth is seen in some cases. Some occur in a site where the permanent tooth is missing. Multiple OCps have been reported [1622, 103A, 2267A] and they may be part of Gardner syndrome [66]. Peripheral lesions developing entirely within the gingival soft tissues are rare [891]. Radiographically, the OCp appears as a collection of tooth-like structures surrounded by a radiolucent zone. Adjacent teeth may be displaced but are never resorbed.

Macroscopy
The specimen consists of a number of tooth-like structures enclosed in a fibrous capsule which is thin if the lesion has matured. In most cases the final diagnosis can be made on the basis of macroscopic examination.

Histopathology
Sections of immature, developing compound odontomas show several dysmorphic tooth germs in a loosely textured connective tissue with cords and islands of odontogenic epithelium. Much of the enamel matrix is preserved in spite of decalcification [2040A].

Prognosis and predictive factors
OCps are treated by local excision. Recurrences have never been reported.
Odontoameloblastoma

Definition
Odontoameloblastoma (OA) combines features of ameloblastoma with those of an odontoma.

ICD-O code
9311/0

Synonyms
Ameloblastic odontoma, odontoblastoma.

Epidemiology
The rare occurrence of this neoplasm precludes definite epidemiologic data (1783). Most OAs have been diagnosed during the first three decades of life (1783).

Localization
This tumour equally affects mandible and maxilla with most of the cases occurring posterior to the canines.

Clinical features / Imaging
Presenting signs may include bone expansion, root resorption, tooth displacement and occasional pain (1288, 1783).
Radiographically, OA appears as a well-defined unilocular or multilocular radiolucent lesion in which varying amounts of radiopaque material may be identified (1783, 2594). Most cases are associated with displaced unerupted teeth.

Macroscopy
Most OAs are unencapsulated. On cut section, the lesion has a multinodular architecture with soft and hard tissue components. The amount of mineralized tissue may appear as large lobulated masses or as rudimentary teeth scattered within the soft tissue.

Histopathology
The epithelial component consists of islands and cords of odontogenic epithelium demonstrating follicular and plexiform patterns, typical of ameloblastoma. In addition to the fibrous stroma, this tumour shows a variable amount of cellular myxoid tissue adjacent to the epithelium, where mineralized dental tissues are formed as in odontomas (1288, 1783, 2594). Isolated small foci of ghost cells may occasionally be found (1783).

Prognosis and predictive factors
Odontoameloblastoma is a locally aggressive neoplasm similar in behaviour and prognosis to conventional ameloblastoma (966, 1288, 1783, 2594).
Calcifying cystic odontogenic tumour

Definition
Calcifying cystic odontogenic tumour (CCOT) is a benign cystic neoplasm of odontogenic origin, characterized by an ameloblastoma-like epithelium with ghost cells that may calcify.

ICD-O code 9301/0

Synonyms
Keratinizing and calcifying odontogenic cyst, Gorlin cyst, calcifying odontogenic cyst

Epidemiology
The CCOT may present as an intraosseous or extraosseous process. The age range varies from 5-92 years without gender predilection [297,298, 1121].

Localization
CCOT shows an equal site distribution for maxilla and mandible. Extraosseous CCOTs usually present in the incisor-cuspid area. Most of the intraosseous CCOTs also are found in the incisor-cuspid area [297].

Clinical features / Imaging
Extraosseous CCOTs are pink to reddish, circumscribed, smooth surfaced, elevated masses, measuring up to 4 cm in diameter. They are usually asymptomatic [298]. Intraosseous CCOTs present as a painless swelling [297]. Radiographs of extraosseous CCOTs may show saucerization and sometimes displacement of adjacent teeth. Intraosseous CCOTs are generally seen as unilocular radiolucencies with a well-circumscribed border. In about 50%, a variable amount of radiopaque material is seen. Root resorption is common, as is root divergence. An associated unerupted tooth is seen in one third of the cases [297,667].

Histopathology
In either variant the cyst wall is lined by a thin ameloblastomatous epithelium with the formation of ghost cells. These ghost cells may calcify [2538A]. Proliferation of odontogenic epithelium in the adjacent connective tissue [1121,2077, 2315] and dysplastic dentin may be observed.

Prognosis and predictive factors
Enucleation is the appropriate treatment for most CCOTs. Recurrence has not been reported for the extraosseous type [298]. A few recurrences have been reported for the intraosseous type [297]. Features of CCOT have been described in a number of other odontogenic tumours [1099,2077].
Dentinogenic ghost cell tumour

**Definition**
A locally invasive neoplasm characterised by ameloblastoma-like islands of epithelial cells in a mature connective tissue stroma. Aberrant keratinization may be found in the form of ghost cells in association with varying amounts of dysplastic dentin.

**ICD-O code** 9302/0

**Synonyms**
Calcifying ghost cell odontogenic tumour, odontogenic ghost cell tumour, epithelial odontogenic ghost cell tumour, dentinoameloblastoma. Formerly dentinogenic ghost cell tumour (DGCT) was considered a solid variant of the calcifying odontogenic cyst (297, 1121, 1498, 2077).

**Epidemiology**
DGCT occurs as an intraosseous and less commonly as an extraosseous variant. The age range is from the second to the ninth decade. The DGCT is somewhat more common in men than in women.

**Localization**
DGCT may occur in any tooth-bearing area of the jaws. There is no preference for maxilla or mandible. The extraosseous variant shows a predilection for the anterior part of the jaws, while the intraosseous variant most often affects the canine to first molar region.

**Clinical features / Imaging**
The tumour is usually asymptomatic [1121, 2527]. The extraosseous variant presents as sessile, sometimes pedunculated, exophytic nodule of the gingival or alveolar mucosa. Many have occurred in edentulous areas. The size varies from 0.5-4.0 cm, but most are between 0.5 and 1 cm. Radiographs will show saucerization of the underlying bone in about 20% of the cases (298, 1553). Teeth in the affected area may be displaced. DGCT is slow growing.

The size of the intraosseous DGCT varies from 1 to more than 10 cm in diameter. It is usually asymptomatic. There may be bony expansion and in some cases resorption of cortical bone with extension into soft tissues. Adjacent teeth may be displaced and mobile. Radiographs show a radiolucent to mixed radiolucent/radiopaque appearance depending on the amount of calcification. The borders are usually well-demarcated. Resorption of adjacent teeth is a common finding, and associated impacted teeth have been described (297, 667).

**Histopathology**
There is no difference between the microscopic features of the intra- and extraosseous variant. The tumour infiltrates the surrounding tissue. Sheets and rounded islands of odontogenic epithelium are seen in a mature connective tissue. The epithelium of the tumour islands resembles that of an ameloblastoma. Mitoses are not seen. Minor cysts may form in the epithelial islands. A characteristic feature is the transformation of the epithelial cells into ghost cells. Individual as well as large islands of ghost cells may be seen. Where basal layer cells are transformed into ghost cells the basement membrane disappears, while ghost cells extrude into the fibrous connective tissue evoking a foreign body reaction. Some ghost cells undergo calcification (723). DGCT forms dysplastic dentin although in minute amounts. Ghost cells may be trapped in the dysplastic dentin, which in some areas may be mineralized. DGCT can be distinguished from ameloblastoma by the presence of large numbers of ghost cells and dysplastic dentin. DGCT may be difficult to distinguish from a multicystic calcifying cystic odontogenic tumour (CCOT). Malignant transformation of a DGCT into an odontogenic ghost cell carcinoma has been described.

**Prognosis and predictive factors**
The intraosseous DGCT may be aggressive with wide local resection recommended, particularly if the tumour is radiologically ill-defined. Enucleation is an appropriate treatment of the extraosseous DGCT; no recurrences have been reported, except in some intraosseous cases, and even malignant transformation has been documented.
Odontogenic fibroma

Definition
The odontogenic fibroma (OF) is a rare neoplasm characterized by varying amounts of inactive-looking odontogenic epithelium embedded in a mature, fibrous stroma.

ICD-O code
9321/0

Synonyms and historical annotation
Controversy exists as to concept and definition [628]. At present the term OF is applied to two histological types of lesions: the epithelium-poor type (formerly termed simple type) and the epithelium-rich type (formerly termed complex or WHO-type).

Epidemiology
Due to lack of uniform definition, data on relative frequency are wide-ranged and inconsistent. When considering the epithelium-rich type, the age range of 15 reported cases was 11-66 years with a mean of 40 years, with a female predominance of 2.8:1 [37,522,609,620, 629, 1206,1618,2045,2270].

Localization
Topographically, two variants can be distinguished: an intraosseous or central type (COF) described here and an extraosseous or peripheral type [530]. From the above-mentioned source of epithelium-rich cases, more were located in the mandible giving a maxilla: mandible ratio of 1:6.5 with most lesions found in the mandibular/premolar area.

Clinical features / Imaging
The epithelium-rich type presents a slow-growing, progressive but painless swelling, often with cortical expansion. In half of cases the tumour appears as a unilocular radiolucent area with well-defined often sclerotic borders. Rarely, the occurrence of calcified material may produce a mixed radiolucent / radiopaque appearance. Larger lesions show scalloping of the margins. Adjacent teeth may be displaced. Some tumours are associated with the crown of an unerupted tooth.

Histopathology
The epithelium-poor type of COF is a non-infiltrating connective tissue lesion resembling a dental follicle. It is minimal-ly cellular with dispersed delicate collagen fibres. A considerable amount of ground substance produces a fibromyxoid quality to the background. Scattered remnants of inactive-looking odontogenic epithelium appear as small irregular islands and cords. Occasionally, variably-formed calcifications occur. The epithelium-rich type of COF is composed of cellular, fibroblastic connective tissue interwoven with less cellular and often vascular areas. Islands or strands of inactive-looking odontogenic epithelium are an integral component; they may be sparse but are often conspicuous. This type shows foci of calcified material considered to be metaplastically produced dysplastic cementum/osteoid/dentin. A well-defined capsule is rare. Subvariants of both histological types of COF have been described [37,628, 1928].

Histogenesis
It has been suggested [863] that the epithelium-poor type of COF is derived from the dental follicle whereas the epithelium-rich type arises from the periodontal ligament. Existence of two types of COF has been challenged [999].

Prognosis and predictive factors
Both types of COF are benign lesions and are cured by local enucleation. Long-term follow-up studies are not available.

Fig. 6.59 Odontogenic fibroma, epithelium-rich type. A Periapical radiograph with a well outlined osteolysis with internal trabeculation. B Collagenous cell rich stroma with numerous strands of odontogenic epithelium without palisading of peripheral cells. C Calcified material encircles an epithelial island.
**Odontogenic myxoma / myxofibroma**

**Definition**
Odontogenic myxoma (OM) is an intraosseous neoplasm characterized by stellate and spindle-shaped cells embedded in an abundant myxoid or mucoid extracellular matrix. When a relatively greater amount of collagen is evident, the term myxofibroma may be used.

**ICD-O code**
9320/0

**Synonym**
Odontogenic fibromyxoma

**Epidemiology**
The frequency of Odontogenic myxoma (OM) varies in different parts of the world between 3-20% of all odontogenic tumours [1925]. In most studies, OM is the third most frequent odontogenic tumour (after odontoma and ameloblastoma). The age range varies from 1-73 years, with a mean age of 30 years. The majority is diagnosed in the 2nd-4th decades. OM is slightly more common in females [1245].

**Localization**
Two-thirds of OMs are located in the mandible. OMs are most common in the molar regions. Maxillary lesions tend to obliterate the maxillary sinuses as an early feature [1245].

**Clinical features / Imaging**
Small OMs are asymptomatic. Large OMs cause painless expansion. Cortical perforation may occur when large. Unilateral sinonasal obliteration may mimic nasal polyposis. Radiographically, OMs appear as unilocular or multilocular radiolucency, sometimes showing a fine “soap bubble” or “honeycomb” appearance occasionally with fine trabeculations. The borders of the tumour are usually well-defined and corticated but can be poorly defined or diffuse. Root displacement occurs, as does root resorption. Larger OMs may present with periosteal reactions. CT may reveal the fine bony septa and allows for anatomic deliniation.

**Macroscopy**
Gross examination reveals a grey-white mass with a typical translucent mucinous appearance. The consistency varies from gelatinous to firm, depending on the amount of collagen present and fine white bands of collagen may be visible on the cut surface.

**Histopathology**
OM is characterized by randomly orientated stellate, spindle-shaped and round cells with long, fine, anastomosing pale or slightly eosinophilic cytoplasmic processes extending from the centrally placed nucleus. Cells are evenly dis-
persed in an abundant mucoid or myxoid stroma that contains only a few fine collagen fibres. Binucleated cells, mild pleomorphism and mitotic figures may occur [1649]. Rests of odontogenic epithelium are not obvious in most lesions and are not required for establishing final diagnosis. Some OMs may permeate into the marrow spaces in a pseudo-malignant pattern. Some OMs have a tendency to produce collagen fibres and are designated myxofibroma. There is no evidence that these more collagenous variants behave differently. Histochemical studies show that the ground substance is rich in acid mucopolysaccharides, primarily hyaluronic acid and, to a lesser degree, chondroitin sulphate. OM is strikingly similar microscopically to myxoid enlarged or 'hyperplastic' dental follicle and the dental papilla of a developing tooth. Misdiagnosis of these entities should be avoided by correlation with the clinical and radiographic features [121,1313,2489]. In maxillary cases, confusion with nasal polyps is a risk. The microscopic differential diagnosis should also include myxoid nerve sheath tumours, chondromyxoid fibroma, low-grade myxoid fibrosarcoma and other myxoid sarcomas.

**Somatic genetics**

A study of 23 cases has shown that odontogenic myxomas are not associated with activating mutations of the Gs alpha gene [239]. Odontogenic myxoma has been reported in a single case of tuberous sclerosis [1019] but is not otherwise associated with Carney complex or any known genetic lesion. In one study karyotypic aberrations have been demonstrated [1969A].

**Prognosis and predictive factors**

The tendency of OM to permeate into marrow spaces makes effective enucleation and curettage difficult. Small lesions have been successfully treated in this way but larger lesions may require complete excision with free margins. Recurrence rates from various studies average about 25% but in spite of this, the prognosis is good. Recurrence usually follows incomplete removal within two years but may occur much later. Death may ensue due to cranial base extension [1969A].
Cementoblastoma

Definition
A cementoblastoma is characterized by the formation of cementum-like tissue in connection with the root of a tooth.

ICD-O code 9273/0

Epidemiology
Just over a hundred cases have been reported. The age range is from 8 up to 44 years, the mean being approximately 20 years (262). There is no distinct gender preference.

Localization
The majority of cementoblastomas are located in the mandible, particularly related to the permanent first molar; association with a primary tooth is exceptional.

Clinical features / Imaging
The most common finding is a painful swelling at the buccal and lingual/palatal aspect of the alveolar ridges. The vitality of the involved tooth remains intact. Lower-lip paresthesia or a pathologic fracture of the mandible are rarely reported (262). Radiographically, the tumour is well-defined and is mainly of a radiopaque or mixed-density, surrounded by a thin radiolucent zone. Root resorption, loss of root outline and obliteration of the periodontal ligament space are common findings.

Macrosopy
The tumour consists of a rounded or nodular mass attached to one or more tooth roots and is surrounded by a grey-to-tan layer of irregular soft tissue (262).

Histopathology
A cementoblastoma consists of dense masses of acellular cementum-like material in a fibrous, sometimes rather vascular stroma that may contain multinucleated cells. The tumour mass blends with the root of a tooth with simultaneous root resorption. In the more mature parts of the tumour, basophilic reversal lines may produce a Paget disease-like pattern. At the periphery sheets of unmineralized tissue may be seen, often being arranged in radiating columns. The differential diagnosis includes osteoblastoma, the only distinctive criterion being the true connection with the surface of the root of a tooth in case of a cementoblastoma (2400). An important differential diagnosis is osteosarcoma. Without radiographs it is difficult to properly diagnose a cementoblastoma. The diagnosis cannot be made on the biopsy alone.

Prognosis and predictive factors
In case of incomplete removal, together with the associated tooth, recurrence is common (262).
**Ossifying fibroma**

**Definition**
Ossifying fibroma (OF) is a well-demarcated lesion composed of fibrocellular tissue and mineralized material of varying appearances. Juvenile trabecular ossifying fibroma (JTOF) and juvenile psammomatoid ossifying fibroma (JPOF) are two histologic variants of ossifying fibroma.

**ICD-O code** 9262/0

**Synonyms**
Cementifying fibroma, cemento-ossifying fibroma, juvenile (active/aggressive) ossifying fibroma

**Epidemiology**
OF most commonly occurs in the 2nd to 4th decades and shows a predilection for females [261]. The mean age of the histological subtypes varies. In patients with JPOF it is about 20 years compared to 35 years in cases of conventional ossifying fibroma [1229]. JTOF has a still lower mean age range (8.5-12 years) [644].

**Localization**
OF is mostly seen in the posterior mandible [261]. JPOF mainly occurs in the bony walls of the paranasal sinuses whereas in the JTOF the maxilla is the site of predilection [644].

**Clinical features / Imaging**
OF causes expansion of the involved bone [261]. Radiographs show a demarcated lesion that may have radiodense as well as radiolucent areas depending on the various contributions of soft and hard tissue components [261].

**Macroscopy**
OFs are well-demarcated, firm lesions without further noteworthy macroscopic features.

**Histopathology**
Ossifying fibroma (OF) is composed of fibrous tissue that may vary in cellularity from areas with closely packed cells to nearly acellular parts within the same lesion. The mineralized component may consist of woven bone, lamellar bone and acellular to poorly cellular basophilic and smoothly contoured deposits thought to be cementum. Due to the presence of this cementum-like material, ossifying fibromas have also been called cemento-ossifying fibroma. However, cementum is defined as a mineralized material covering the surface of the roots of the teeth and outside this location, its distinction from bone is equivocal and without clinical relevance [261,2401]. OF may be confused with fibrous dysplasia. The most important distinguishing feature is the presence of demarcation and/or encapsulation in OF as opposed to the merging with its surroundings as shown by fibrous dysplasia. In addition, the variation in cellularity as well as in
appearances of mineralized material serves to distinguish OF from fibrous dysplasia \cite{2401,2699}. Distinction between OF and osseous dysplasia on histologic grounds only may be problematic as both entities share the variation in stromal cellularity and appearances of mineralized material. Clinical presentation and radiographic appearance may be decisive (see osseous dysplasia).

**Juvenile trabecular ossifying fibroma (JTOF)** consists of cell-rich fibrous tissue containing bands of cellular osteoid without osteoblastic rimming together with slender trabeculae of immature bone containing coarse lacunae with plump osteocytes and are lined by a dense rim of enlarged osteoblasts. Sometimes these trabeculae may anastomose to form a lattice. Mitoses are present, especially in the cell-rich areas. Additional but less typical features are multinucleated giant cells, pseudocystic stromal degeneration, and haemorrhages \cite{644,2051,2406}.

**Juvenile psammomatoid ossifying fibroma (JPOF)** is characterized by a fibroblastic stroma containing small ossicles resembling psammoma bodies. The stroma varies from being loose and fibroblastic to intensely cellular with minimal intervening collagen. The mineralized material consists of spherical or curved ossicles that are acellular or show sparsely distributed cells. These ossicles should not be confused with the cementum-like deposits that are present in conventional OF. These particles have a smooth contour with sometimes a radiating fringe of collagen fibers, whereas the ossicles in JPOF have a thick irregular collagenous rim that may attain such a size that it includes multiple ossicles. The ossicles themselves may also fuse to form trabeculae showing reversal lines. Sometimes, JPOF contains deeply basophilic concentrically lamellated particles as well as irregular thread-like or thorn-like calcified strands in a hyalinized background. Other features such as trabeculae of woven bone as well as lamellar bone, pseudocystic stromal degeneration and haemorrhages result in areas similar to an aneurysmal bone cyst. Multinucleate giant cells, and mitotic figures may be present, but are not specific for this variant of ossifying fibroma. JPOF has to be distinguished from extra-cranial meningioma with psammoma bodies, which demonstrates EMA positivity. Moreover, the psammomatoid ossicles in JPOF are clearly different from the acellular spherical true psammoma bodies \cite{2599}.

**Histogenesis**

OFs originate from the periodontal ligament \cite{2861}.

**Somatic genetics**

The following chromosomal abnormalities have been observed in OF: one case with three reciprocal translocations with the karyotype 46,XY,t(1;18)(q21;q21.3), t(3;10)(P13;q22),t(6;11)(p22;p15), one case with alterations affecting the short arms of chromosomes X, 2, and 7 and 3 cases with identical chromosomal breakpoints occurring at bands Xq26 and 2q33, with an identical t(X;2)(q26;q33) reciprocal translocation in 2 cases and an interstitial insertion of bands 2q24.2q33 into Xq26 in the third case \cite{528,919,2254}.

**Prognosis and predictive factors**

OFs continue to enlarge when left untreated. Therefore, they should be removed completely.
Fibrous dysplasia

Definition
Fibrous dysplasia (FD) is a genetically-based sporadic disease of bone that may affect single or multiple bones (monostotic: MFD, polyostotic: PFD). FD occurring in multiple adjacent craniofacial bones is regarded as monostotic (craniofacial FD). FD may be part of the McCune-Albright syndrome (MAS).

Epidemiology
The monostotic form (MFD) is equally distributed in both genders and ethnic groups and is six times more common than the polyostotic: PFD [775]. PFD is more frequent in females (F/M ratio, 3:1). MFD and PFD are mainly diagnosed in children and young adults. However, in the 3% of all PFD-cases that occur in the setting of McCune-Albright syndrome, the disease may manifest in infants [775, 1018].

Etiology
Mutations in the gene (GNAS I) encoding for the α-subunit of a signal transducing G-protein (Gs-α) lead to increased c-AMP production affecting proliferation and differentiation of preosteoblasts [472,1633,2175].

Localization
In the jaws, FD occurs more often in the maxilla than in the mandible, and may involve adjacent bones like the zygoma or the sphenoid [93,2710]. The long tubular bones, especially the femur, followed by the flat bones of the jaws, the skull (base of skull prior to neurocranium), and the ribs, are the most frequently affected sites in the skeleton [2067].

Clinical features
Signs and symptoms
Complaints usually consist of painless swelling often leading to facial asymmetry, occasionally accompanied by irregular café-au-lait spots. Maxillary and mandibular involvement may lead to displacement of teeth, malocclusion and, rarely, root resorption [1759]. In FD affecting the paranasal sinuses nasal obstruction may occur [2321]. Lesions extending to the orbit may cause visual impairment, while temporal bone lesions may produce hearing loss. Sometimes facial pain, headaches or facial numbness develops [140,281,1586]. Children presenting with FD, especially PFD, and irregular café-au-lait spots should be carefully examined for MAS [1002]. An elevation of alkaline phosphatase, even in the absence of fractures and unrelated to the extent of the disease has been noted in up to one third of FD patients [1018].

Imaging
Three different radiographic patterns of FD involving the maxillofacial skeleton have been described: cystic (radiolucent or lytic; early lesions), sclerotic (mid-phase lesions), and mixed radiolucent/radiopaque (pagetoid: late lesions) comprising 21%, 23% and 56%, respectively [281,820,1759]. Asymmetric homogeneously radiodense opacities with “ground glass”-appearance that blend into normal bone, thin cortices and bone expansion are highly characteristic for FD and best seen on CT scans on bone windows [520]. In the jaws, superior displacement of the mandibular canal, narrowing of the periodontal ligament space and effacing of the lamina dura are findings suggestive of FD [520,1759,2024, 2572]. On MRI, signals are intermediate.

Fig. 6.76 Fibrous dysplasia. Polyostotic fibrous dysplasia (mandible, base of skull) with extreme deformity of the mandible. Frontal (coronal) CT scan, bone window.

Fig. 6.75 Fibrous dysplasia of the jaw. Radiograph showing expansile osteolysis with irregular opacities of the mandible, extending into the ascending ramus up to the mandibular condyle. Partially eroded lamina dura of affected teeth.
on T1 and proton-weighted images, and heterogeneous hypointense on T2, showing a moderate to marked signal enhancement following Gd-DPTA administration [1586,1759].

**Histopathology**
FD consists of cellular fibrous tissue with spindle shaped cells and immature, isolated trabeculae of woven bone generally without rimming of osteoblasts [2401, 2710]. Characteristically, bundles of collagen fibres oriented perpendicular to the bone surface, compatible with Sharpey-fibers, can be demonstrated [1018,2067,2176]. Osteoid seams are present and best visualized on undecalcified sections [210,2067]. In long standing lesions some osteoblastic rimming and “maturation” to lamellar bone may occur. This may result in parallel ordered bony trabeculae [2379,2401]. Cartilaginous foci in FD of the jaws or skull have not been documented.

**Genetics**
Activating mutations in the GNAS1 gene, coding for the α-subunit of the stimulatory G-Protein have been proven in MFD, PFD as well as in MAS [472,2175]. In eight of eleven cases clonal chromosomal aberrations have been described, including both structural and numeric changes. Repeated chromosomal changes have only been documented so far for trisomy 2 and rearrangements of 12p13, in three cases each [527]. These findings may suggest that FD is a neoplastic process [472,527,775].

**Prognosis and predictive factors**
In most cases of FD the lesions seem to stabilize with skeletal maturation. Surgical interventions may be necessary for functional reasons or severe disfigurement [2710]. Very rarely, sarcoma development, predominantly osteosarcoma, has been reported preferentially in craniofacial bones and even in the absence of prior irradiation [220,2213, 2522,2835].
Osseous dysplasias

Definition
Osseous dysplasias (ODs) are idiopathic processes located in the periapical region of the tooth-bearing jaw areas, characterized by a replacement of normal bone by fibrous tissue and metaplastic bone.

Synonyms
Periapical cemental dysplasia, periapical osseous dysplasia, focal cemento-osseous dysplasia, periapical cementoma.

Epidemiology
OD has a predilection for middle-aged black females [261].

Localization
OD is confined to the tooth-bearing part of the jaws.

Clinical features / Imaging
The condition occurs in various clinical forms that bear different names. When occurring in the anterior mandible and involving only a few adjacent teeth, it is called periapical osseous dysplasia. A similar limited lesion occurring in a posterior jaw quadrant is known as focal osseous dysplasia, formerly called focal cemento-osseous dysplasia [2502]. Two other types of OD are more extensive, occurring bilaterally in the mandible or even involving all 4 jaw quadrants. The first is known as florid osseous dysplasia [261,2864]. This type of OD mainly occurs in middle-aged black females. The second occurs at young age and causes considerable jaw expansion. This OD type is called familial gigantiform cementoma; it shows an autosomal dominant inheritance with variable expression but sporadic cases without a history of familial involvement have been reported [7, 2864]. Periapical and focal OD usually are incidental radiographic findings. The involved teeth remain vital. Florid OD may give rise to symptoms in cases of concomitant infection. Jaw expansion is not a feature of ODs with the exception of the familial gigantiform cementoma and is rarely seen in florid OD.

Histopathology
All types of OD consist of cellular fibrous tissue, woven as well as lamellar bone and masses of cementum-like material. There is no capsule. The hard tissue component in most cases does not fuse with the root surface of the involved teeth, but may merge with the surrounding bone. Secondary inflammatory changes may occur, especially with the florid OD and the familial gigantiform cementoma. OD resembles ossifying fibroma (OF) histologically. Clinical and radiographic information is needed to make the distinction [2485]. OD may be also confused with fibrous dysplasia (FD). However, the variation in appearances of mineralized material distinguishes both lesions, fibrous dysplasia almost exclusively consisting of woven bone [2401].

Histogenesis
Osseous dysplasia is considered to originate from the periodontal ligament [2861].

Prognosis and predictive factors
The various forms of osseous dysplasia do not require treatment unless complications occur such as infection of sclerotic bone masses as may be encountered in florid OD or facial deformity as may be seen in familial gigantiform cementoma.
Central giant cell lesion

Definition
Central giant cell lesion (CGCL) is a localized benign but sometimes aggressive osteolytic proliferation consisting of fibrous tissue with haemorrhage and haemosiderin deposits, presence of osteoclast-like giant cells and reactive bone formation.

Synonyms
Central giant cell granuloma, reparative giant cell granuloma.

Epidemiology
CGCL is found in all age groups, however, most cases are diagnosed in patients under 30 years of age, with an incidence rate of 1.1/million population / year [559]. In contrast to giant cell tumour of bone, about 1/3 of patients are younger than 20 years [1239]. Women are more often affected than men (1.5-2:1) [1244].

Localization
The mandible is more often involved than the maxilla [1244,2782]. Molar and premolar areas are more often affected than the anterior parts or the ascending ramus [2460]. Involvement of the condyle or maxillary sinus is rare [11, 1244]. Multifocal giant cell lesions do occur. They may be sporadic and unrelated to other conditions (eg. hyperparathyroidism, cherubism, Noonan syndrome) [2414,2663].

Clinical features / Imaging
Most cases present as asymptomatic incidental findings. Some, however, present with pain or paraesthesia, swellings, or loosening of teeth. Nasal obstruction may occur [1239,2782]. CGCL are expansile, radiolucent and often multiloculated lesions, rarely mixed with opacities, with scalloped and mostly well-defined but non-corticated borders. With increasing size, multilocularity is more often noticed [2460]. Disappearance of the lamina dura, root resorption or, more often, tooth displacement are additional findings [452,967,1244,2460,2782]. Intralesionary wavy bony septa are characteristic [967]. Periapical localization may mimic periapical granuloma [524].

Histopathology
The lesion consists of spindle-shaped fibroblastic or myofibroblastic cells [643,1920], loosely arranged in a fibrous, sometimes fibromyxoid, vascularized tissue with haemorrhagic areas, haemosiderin deposits, macrophages, lymphocytes, granulocytes and, rarely, plasma cells. Especially in the haemorrhagic areas, evenly dispersed or small clusters of osteoclast-like giant cells are found [84,452,771]. In addition, traversing collagen bundles are present, often accompanied by metaplastic bone formation, giving the lesion a somewhat lobular appearance. Mitoses are frequently found [1239,2398,2839]. Since brown tumour of hyperparathyroidism is morphologically indistinguishable from CGCL, determination of parathormone levels may be indicated, especially in elderly patients or when multifocal lesions are present [1276,1645,2694].

Genetics
In one case of a peripheral CGCL of the distal phalanx a cytogenetic study has been performed revealing a translocation involving the X-chromosome and chromosome 4 [t(X;4)(q22;q31.3)] [310].

Prognosis and predictive factors
Histological findings are not predictive of biological behaviour. The treatment of CGCL is careful enucleation. In case of recurrences, more extensive surgery should be considered. Administration of calcitonin (intranasal or s.c), [558,1015,1921] or glucocorticoids (intralesionary) has proven effective in some cases [333,1394]. More recently, antiangiogenic therapy with interferon alpha has been successfully applied [1241].
Cherubism

Definition
Cherubism is an autosomal dominant inherited disease that is characterized by a symmetrical distension of the jaws, often leading to a typical facial expression. The histology is indistinguishable from central giant cell lesion.

Epidemiology
Cherubism is a familial disease affecting 100% of males and up to 70% of females. Sporadic cases do occur (589,1237,2023). Generally, diagnosis is made in early childhood (14 months to 4 years) or, in milder forms, in pre-adolescence. With increasing age, especially after cessation of bone growth, the lesions regress (2608).

Localization
All four quadrants of the jaws can be involved. Usually the mandible is affected more extensively, starting at the angle at the time of permanent molar eruption. The process may extend into the ascending ramus without affecting the condyle, and the mandibular body. In the maxilla, both tuberosities are affected initially followed by involvement of the anterior and inferior portions of the orbits (76,479).

Clinical features / Imaging
Symmetrical swellings and an indolent clinical course are characteristic. Bilateral maxillary enlargement may lead to retraction of the facial skin including the lower eyelids, resulting in scleral exposure and the typical "looking toward Heaven" appearance (cherubs on Renaissance paintings) (479,1236). Other consequences are tooth displacement and delay in tooth eruption, loosening of teeth, speech alterations and visual impairment. In addition, cervical lymphadenopathy is noticed (24,76,1287,2608). Affected bones are expanded by bilateral, well-delineated multilocular radiolucencies with a "soap bubble" appearance. The cortices may become thinned and focally perforated. With advancing age, the initially fibrous tissue is replaced by bony structures, leading to sclerosis (76). The diagnosis is supported by clinical presentation (bilateral enlargement of the jaws) and the typical radiological findings on panoramic or lateral views, or CT-scans (1105).

Histopathology
Initially, fibrous tissue and giant cells resembling osteoclasts are present, giving an impression that may be almost indistinguishable from central giant cell lesion (1287,2840). Additional features are haemosiderin deposits and stromal fibrosis (1237). Although a rare finding, perivascular cuff-like collagen deposits are regarded as characteristic for cherubism (990). Although the histology is not specific, the combination of clinical appearance, radiology and central giant-cell lesion-like histology is diagnostic.

Genetics
Cherubism is an autosomal dominant familial disease and has been mapped to chromosome 4p16.3 (1621,2608). Through analysis of 12 families, the mutated gene was recently identified as SH3BP2, coding for a c-Abl-binding protein. However, since the mutation was not detected in three other families, mutations in genes different from SH3BP2 cannot be ruled out (2650).

Prognosis and predictive factors
With time, especially after puberty, the lesions regress (1286). Before puberty, surgery should be carried out only in cases of severe functional disturbances (1362).
Aneurysmal bone cyst

Definition
Aneurysmal bone cyst (ABC) is an expansile osteolytic lesion often multilocular, with blood filled spaces separated by fibrous septa containing osteoclast-type giant cells and reactive bone.

Epidemiology
ABC has an incidence of 0.014/100,000 and occurs preferentially in patients below the age of 30 years with a peak in the second decade [1464]. Epidemiologic data for the jaws are lacking. However, ABC in the jaws is rare encompassing 1-3% of all ABCs [1239].

Etiology
ABC may arise primarily or as a secondary event in another bone lesion, e.g. giant cell lesion or fibrous dysplasia [2483, 2802]. Most lesions are believed to be reactive, however, an association with trauma is unlikely, especially for maxillary ones [150]. Cytogenetic data provide evidence that at least some ABCs are neoplastic.

Localization
ABC is more often found in the mandible [1303] with a predominance for the posterior regions including the ascending ramus [1246]. The condyle is only exceptionally affected [1787]. A more uniform distribution is noted in the maxilla [150].

Clinical features
ABC may present with marked swelling and few symptoms. Malocclusion, tooth displacement and loosening may develop. Teeth remain vital [150, 2482]. Root resorption may be present [1246]. Orbital involvement may lead to exophthalmos and diplopia. Nasal obstruction or bleeding is rare [150].

Imaging
Unilocular or multilocular radiolucent lesions are seen. In up to 10% a mixed radiopaque-radiolucent pattern can be found. Borders are well delineated but perforations of the cortices may be present and extensions into the soft tissues do occur. On cross sectional imaging studies internal septa are visible. Fluid-fluid levels produced by sedimentation of blood cells in lesional cavities are particularly seen on MRI (gradient echo sequences or T2 weighted images) and are very characteristic [78, 1246, 2160].

Histopathology
ABC is haemorrhagic, multilocular and well circumscribed. The blood-filled cavities are lined by macrophages and not by endothelial cells [39]. They represent pseudocysts since there is no epithelial lining. The septa are composed of inconspicuous fibroblasts, osteoclast-like giant cells and reactive bone or irregular osteoid are distributed parallel to the septal lining. Haemosiderin deposits are also present. Mitoses are frequently found, however, atypical forms are not seen. Necrosis may be present [1303, 2802].

Genetics
A 17p-rearrangement is a constant finding in ABC in the extracranial skeleton, most often presenting as balanced translocation with 16q but other chromosomes may also be involved [147, 1072, 1894, 1977, 2281, 2832]. One case originating in the nose revealed a (6;17)(p21;p13) translocation [2797]. Since the metaphases of the non-involved chromosomes seem to be normal the translocations may be regarded as resulting from acquired aberrations providing evidence, that at least some ABCs are clonal proliferations [775]. Familial cases have been reported [598, 943, 2692].

Prognosis and predictive factors
ABC can be treated with curettage [1249]. Soft tissue extension increases the risk of recurrence. Embolization has also been applied successfully [956].
Simple bone cyst

Definition
Simple bone cyst (SBC) is an intraosseous pseudocyst devoid of an epithelial lining, either empty or filled with serous or sanguinous fluid.

Synonyms
Solitary bone cyst, traumatic bone cyst, haemorrhagic bone cyst, haemorrhagic cyst, unicameral bone cyst, idiopathic bone cavity

Epidemiology
SBC is most commonly observed during the second decade of life [184]. In contrast to the long bones (male/female ratio of 2:3:1), in the jaws there is no gender predilection [1392].

Etiology
The etiology is unknown [1752,2341].

Localization
The mandible [1239] is almost exclusively affected [1239,1392]. Most SBCs are located in the anterior part of the mandible [1392]. Cases with bilateral lesions have been described [1990].

Clinical features
In general, SBC is discovered incidentally. Complaints are tenderness or mild pain. Displacement of teeth or pathological fractures are very unusual [2341]. Teeth are usually vital [1006]. A history of trauma is rarely reported [1665].

Imaging
Usually, SBC is radiolucent and unilocular with no or only slight expansion of bone and cortical thinning [1665]. Superior margins extend between the roots of teeth and are characteristically scalloped and corticated [2272]. Root resorption is an unusual finding. Intracavitary fluid has been visualized on T2 weighted MRI images [690]. A final diagnosis is aided by finding an empty cavity at surgical exploration.

Histopathology
The lining of the cavity is made up of connective tissue covering the underlying bone with a membrane-like layer [184,1239]. Rarely, a thickened myxofibromatous wall is seen [1665,2224]. Small amounts of new bone formation and collagen deposits may be present, often described as appearing fibrin- or cementum-like. In addition, scattered giant cells and haemosiderin are also found [1006].

Genetics
Only two genetically analyzed cases, localized in the long bones, have been reported so far. Complex clonal structural rearrangements of chromosome 4, 6, 8, 16, 21, and both 12 were demonstrated [2686]. In addition, a translocation t(16;20)(p11.2;q13) present as the sole abnormality has been described [2166].

Prognosis and predictive factors
Usually bone healing is completed within a year after surgical exploration. However, persistence following curettage may occur requiring additional treatment [789,1392].

Fig. 6.88 SBC. A Centrally located unicameral osteolysis with sclerotic margins extending between the roots of adjacent teeth. Panoramic radiograph. B Expansion of mandible with thinning of corticalis.

Fig. 6.89 Simple bone cyst (SBC). A Membrane-like fibrous covering of the cavity overlying bone. B Thinned corticalis covered by a membranous layer and fibroblasts on top. Focal deposition of collagen (top right).

Fig. 6.90 SBC. Thinned cortical bone covered by fibrous septa.
CHAPTER 7

Tumours of the Ear

Tumours are unusual in the ear. In the external ear most of the neoplasms are those of the covering skin. Only the ceruminous glands are peculiar to the external ear, but ceruminous tumours are rare. The underlying bone contributes some swellings and neoplasms to this area. The most common tumour in the middle ear is the adenoma, which arises from low-mitotic cuboidal epithelium that may become neoplastic. The inner ear is composed of a specific inert bone, a virtually non-mitotic sensory area and nerves. Tumours that are derived from Schwann cells are the only frequent neoplasms of the inner ear, indeed of the whole temporal bone. Diagnosis of ear tumours presents a peculiar difficulty in that the whole structure is often encased in dense bone. Although modern imaging techniques have helped greatly to identify tumours and tumour-like lesions of the ear, there is still a need for autopsy studies in this area.
### WHO histological classification of tumours of the ear

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<th>Papillary tumours</th>
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<tbody>
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<td><strong>Aggressive papillary tumour</strong></td>
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<tr>
<td>Adenoma</td>
<td>8420/0</td>
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<tr>
<td>Chondroid syringoma</td>
<td>8940/0</td>
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<tr>
<td>Syringocystadenoma papilliferum</td>
<td>8406/0</td>
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<tr>
<td>Cylindroma</td>
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<td><strong>Inverted papilloma</strong></td>
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<td>Angiolymphoid hyperplasia with eosinophilia</td>
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<tr>
<td><strong>Tumours of the middle ear</strong></td>
<td><strong>Squamous cell carcinoma</strong></td>
</tr>
<tr>
<td>Adenoma of the middle ear</td>
<td>8140/0</td>
</tr>
<tr>
<td><strong>Tumours of the inner ear</strong></td>
<td><strong>Meningioma</strong></td>
</tr>
<tr>
<td><strong>Papillary tumours</strong></td>
<td>8260/1</td>
</tr>
<tr>
<td><strong>Schneiderian papilloma</strong></td>
<td>8121/0</td>
</tr>
<tr>
<td><strong>Inverted papilloma</strong></td>
<td>8121/1</td>
</tr>
<tr>
<td><strong>Squamous cell carcinoma</strong></td>
<td>8070/3</td>
</tr>
<tr>
<td><strong>Meningioma</strong></td>
<td>9530/0</td>
</tr>
<tr>
<td><strong>Tumours of the inner ear</strong></td>
<td><strong>Vestibular schwannoma</strong></td>
</tr>
<tr>
<td><strong>Lipoma of the internal auditory canal</strong></td>
<td>9560/0</td>
</tr>
<tr>
<td><strong>Haemangioma</strong></td>
<td>8850/0</td>
</tr>
<tr>
<td><strong>Endolymphatic sac tumour</strong></td>
<td>9120/0</td>
</tr>
<tr>
<td><strong>Haematolymphoid tumours</strong></td>
<td><strong>9823/3</strong></td>
</tr>
<tr>
<td>B-cell chronic lymphocytic leukaemia / small</td>
<td><strong>Langerhans cell histiocytosis</strong></td>
</tr>
<tr>
<td>lymphocytic lymphoma</td>
<td>9670/3</td>
</tr>
<tr>
<td><strong>Secondary tumours</strong></td>
<td>9751/1</td>
</tr>
</tbody>
</table>

1. Morphology code of the International Classification of Diseases for Oncology (ICD-0) (821) and the Systematized Nomenclature of Medicine (http://snomed.org). Behaviour is coded /0 for benign tumours, /3 for malignant tumours, and /1 for borderline or uncertain behaviour.
Ceruminous gland neoplasms of external auditory canal and cylindroma

Definition
External ear neoplasms derived from ceruminous glands are very uncommon and can be benign or malignant. Only the adenoma (ceruminoma) can be categorized as being derived specifically from ceruminous glands. Syringocystadenoma papilliferum and adenoid cystic carcinoma arising in this region can sometimes manifest an origin from ceruminous glands. These tumours are either benign or malignant.

Localization
The expected site of origin is in the superficial part of the external canal.

Clinical features
The symptoms of this lesion, like other external ear canal lesions, are conductive hearing loss and discharge. Pain and facial nerve palsy are clinical predictors of malignancy.

Epidemiology
The benign and malignant tumours occur with equal frequency in men and women with a mean age of 49 years (range 26-89 years) (569,1478,1589).

Adenoma of ceruminous glands

ICD-O code 8420/0

Macroscopy
Gross appearances are those of a non-ulcerating superficial grey mass up to 4 cm in diameter, which is covered by skin.

Histopathology
Microscopically this neoplasm lacks a capsule. It is composed of regular oxyphil glands often with intraluminal projections. The glandular epithelium is bilayered. The outer myoepithelial layer may not be obvious in all parts of the neoplasm. In some ceruminomas, acid-fast fluorescent ceroid pigment may be found which is similar to that seen in normal ceruminal glands (2778).

Electron microscopy. One case of ceruminous gland adenoma showed apocrine caps, microvilli, cell junctions, secretory granules, vacuoles, lipid droplets and siderosomes, the characteristic ultrastructural features of apocrine glands (2260).

Chondroid syringoma

Definition
Benign tumour similar to the pleomorphic adenoma of salivary glands.

ICD-O code 8940/0

Synonym
Pleomorphic adenoma or mixed tumour.

Histopathology
Cartilage, myoepithelial and adenomatous structures are features of this neoplasm.

Syringocystadenoma papilliferum

Definition
Benign adnexal tumour with features similar to those seen at other sites.

Fig. 7.1 Ceruminous adenoma. Keratinized squamous epithelium overlies a circumscribed but unencapsulated neoplastic proliferation of ceruminous glands. Note glandular and small cystic profiles.

Fig. 7.2 Ceruminous adenoma. Stratification of the nuclei with moderate nuclear pleomorphism and a mitotic figure (upper left); Abundant eosinophilic-granular cytoplasm in the luminal cells which show focal decapitation secretion (upper right); glandular structures separated by fibrous connective tissue (lower left); inner luminal secretory cells subtended by basal myoepithelial cells demonstrate the dual cell population (lower right).
**Tumours of the ear**

ICD-O code 8406/0

**Synonym**
Hidradenoma papilliferum

**Epidemiology and localization**
Syringocystadenoma papilliferum is seen in children or young adults usually on the scalp or face. Occasionally it occurs in the ear canal.

**Histopathology**
Cystic invagination from surface epithelium. Projecting into the lumen are papillae covered by bilayered apocrine glandular epithelium which may show decapitation secretion typical of ceruminous glands.

**Cylindroma**

**Definition**
Cylindroma is a benign tumour arising from the epidermal adnexae, whether apocrine- or eccrine-derived is not conclusively known.

ICD-O code 8200/0

**Synonym**
Turban tumour

**Localization**
In the external ear the lesion may be present on the pinna or in the external canal. In these situations it may be part of a multiple “turban tumour” presentation of this neoplasm on the scalp.

**Histopathology**
It is composed histologically of rounded masses of small, darkly staining cells which fit together in a jigsaw-like pattern and are surrounded by pink-staining hyaline material. Extracellular hyaline globules are often present in the cellular masses. Larger cells with vesicular nuclei are also seen [2804]. In contrast to primary adenoid cystic carcinoma, cylindroma in the external canal does not have a cribriform structure, but does have larger cells with vesicular nuclei.

---

**Fig. 7.3** Ceruminous adenoma. A Yellow-brown “ceroid” lipofuscin-like material is seen in the cytoplasm of ceruminous cells, a feature seen in modified ceruminous sweat glands and in ceruminous adenomas. B Glandular structures show ceruminous decapitation secretion in the luminal cells subtended by a prominent, well-defined myoepithelial cell layer (left). The myoepithelial cell nuclei are accentuated with a p63 immunoreaction (right). C Differential immunohistochemical staining highlights the luminal cells (CK7, left) while CK5/6 accentuated the basal cells (right).

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**Fig. 7.4** Syringocystadenoma papilliferum of external ear canal. Note papillae lined by bilayered glandular epithelium projecting into a cystic lumen. There is also a prominent epidermoid cyst.

**Fig. 7.5** Cylindroma of pinna with multiple spherical lesions on pinna, face and temporal region. From L. Michaels & H. Hellquist (1711).

**Fig. 7.6** Cylindroma of pinna showing jigsaw-like pattern of cell groups, surrounded by hyaline basement membranes.
Ceruminous gland neoplasms of external auditory canal

**Malignant tumours of ceruminous glands**

**Definition**
An infiltrating neoplasm derived from ceruminous glands.

**ICD-O codes**
- Adenocarcinoma 8420/3
- Adenoid cystic carcinoma 8200/3
- Mucoepidermoid carcinoma 8430/3

**Localization**
Superficial part of the external ear canal. Origin from the adjacent parotid salivary gland should be excluded.

**Histopathology**
- **Low and high-grade adenocarcinoma**
  These neoplasms possess a glandular structure with evidence of apocrine differentiation and infiltration. Low-grade tumours show loss of a myoepithelial layer and infiltration. The cells of high-grade tumours are markedly atypical with increased mitotic activity and widespread invasion.

- **Adenoid cystic carcinoma**
The microscopic features of these tumours are indistinguishable from those arising in salivary glands. They characteristically widely infiltrate adjacent tissues and invade nerve sheaths.

- **Mucoepidermoid carcinoma**
The tumours arising in this location are usually low-grade and the microscopic features are similar to those arising in salivary glands.

**Prognosis and predictive factors**
Recurrence often complicates surgical removal of high-grade tumours. Death due to involvement of local vital structures and metastases has been reported. Relentless, although often delayed recurrence and eventual bloodstream metastasis, particularly to the lungs is likewise a feature of adenoid cystic carcinoma.
**Squamous cell carcinoma of the external ear**

**Definition**
This malignant tumour of stratified squamous epithelium arises from the normal epidermal covering of the external canal of the pinna.

**ICD-O code**
8070/3

**Synonyms**
Epidermoid carcinoma, squamous carcinoma

**Epidemiology**
The average age at diagnosis is 65-70 years for the pinna lesions and there is a male predominance. The age at presentation is 52-55 years for the external canal tumours which show a female predominance [1226].

**Etiology**
Actinic overexposure and frostbite have been suggested as causes of the pinna lesion. The canal tumours have been linked with the same tumour type in the middle ear as possibly resulting from prolonged chronic inflammation. It is possible, however, that the clinical impression of chronic inflammation has been mistaken, the patients’ symptoms being the result of an occult squamous cell carcinoma.

**Localization**
The majority of squamous cell carcinomas of the external ear arise on the pinna; a lesser number arise in the external canal. The external ear sites of involvement in the pinna in a study of 52 patients are shown in Table 7.1. Rarely there is bilateral external ear involvement [2807].

**Clinical features**
The pinna lesions being in an exposed position are identified early. A serious problem with the canal lesions is the delay in diagnosis because of the minimal symptoms that may be present. Pain, hearing loss and drainage of blood or pus are the main features in that group. A plaque-like or even polypoid mass may be felt or even seen.

**Macroscopy**
Squamous cell carcinomas arising on the pinna grossly resemble those seen elsewhere on the skin. The appearances of the canal lesions are those of a mass, sometimes warty, occluding the lumen and invading deeply into the surrounding tissues. There may be dissolution of the tympanic membrane with invasion of the middle ear. Occasionally, the well-differentiated lesions may not be detected clinically until well advanced.

**Tumour spread and staging**
The TNM staging for skin does not seem applicable at this site because of the presence of cartilage invasion.

**Histopathology**
Epidermoid carcinoma of the external ear usually shows significant degrees of keratinization. Those showing a spindle cell morphology must be differentiated from melanomas and soft tissue tumours.

**Table 7.1 Sites of involvement of squamous cell carcinoma of the pinna in 52 patients (2336).**

<table>
<thead>
<tr>
<th>Site</th>
<th>Number of Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Helix</td>
<td>27</td>
</tr>
<tr>
<td>Posterior auricle</td>
<td>11</td>
</tr>
<tr>
<td>Antihelix</td>
<td>6</td>
</tr>
<tr>
<td>Triangular fossa</td>
<td>3</td>
</tr>
<tr>
<td>Concha</td>
<td>3</td>
</tr>
<tr>
<td>Lobule</td>
<td>2</td>
</tr>
</tbody>
</table>

**Precursor lesions**
Actinic keratosis may precede squamous cell carcinoma.

**Prognosis and predictive factors**
Squamous cell carcinoma of the pinna is an aggressive disease with a high propensity for local recurrence. Tumours confined to the external ear usually have a good outlook after surgical therapy. The outcome of the disease following surgical excision is related to the clinical stage at presentation, the higher the stage the worse the outcome [1915]. Metastatic spread of squamous carcinoma of the pinna and external auditory meatus to lymph nodes is unusual. Lesions arising in the canal have a worse prognosis because of the late diagnosis and invasion of adjacent structures.

Fig. 7.8 Squamous cell carcinoma of the pinna forming a large mass with central ulceration.
Embryonal rhabdomyosarcoma

Rhabdomyosarcoma and its variants have been comprehensively discussed in the WHO Classification of Tumours of Soft Tissue and Bone (775). This section focuses on its occurrence as a primary tumour in the external ear canal.

**Definition**
A primitive malignant tumour with phenotypic and biological features of embryonic skeletal muscle.

**ICD-O code**
8900/3

**Synonyms**
Myosarcoma, embryonal sarcoma, botryoid sarcoma.

**Epidemiology**
Rhabdomyosarcoma is rare in any part of the body. There is a distinct group arising in the head and neck of children, often very young, with a predilection for the palate, middle ear and orbit.

**Localization**
Most of the tumours arise in the middle ear with extension into the external canal as an “aural polyp”.

**Clinical features**
Embryonal rhabdomyosarcoma should be excluded in any child presenting with a polyp in the external ear canal. Advanced cases may present with aural discharge, facial weakness and swelling in the region of the ear (1116). Extensive destruction of the bone at the base of the skull, especially the petrous bone has been described.

**Histopathology**
Only the embryonal subtype of rhabdomyosarcoma is recognized as occurring at this site. The characteristics of this polypoid tumour are those of rhabdomyoblasts and primitive mesenchymal cells showing a variable degree of skeletal muscle differentiation loosely arranged but with condensation beneath the epithelium (cambium layer). Yolk sac tumour has been described as a polypoid tumour presenting in the external ear canal. However, this is histologically distinct, being composed of small round blue cells arranged in a vacuolated pattern with formation of Schiller-Duval bodies and expressing alpha fetoprotein (833). A detailed description of embryonal rhabdomyosarcoma including immunophenotype is given in the WHO Classification of Tumours of Soft Tissue and Bone (775).

**Histogenesis**
Although it is suggested that this tumour arises from striated muscle fibres in the middle ear, it seems more likely that the origin is from undifferentiated mesenchymal cells.

**Genetics**
Mutations in a region mapped to the short arm of chromosome 11 (11p15) have been associated with most embryonal rhabdomyosarcomas. Several genes have been mapped to this site. Complex structural and numerical chromosomal rearrangements have been associated with embryonal rhabdomyosarcoma. These are discussed in detail in the WHO Bone and Soft tissue book.

**Prognosis and predictive factors**
Modern chemotherapeutic schedules have dramatically improved the outcome for children with this tumour.

---

**Fig. 7.9**

A central area of necrosis is surrounded by “primitive cells” with a very high nuclear to cytoplasmic ratio. The neoplasm is separated from the surface. This polypoid tumour has a “Grenz-Zone” between the neoplastic cells and the mucosal surface. The malignant cells have abundant eosinophilic cytoplasm.
**Fibrous dysplasia**

**Definition**
Fibrous dysplasia (FD) is a benign localised intramedullary proliferation of trabecular woven bone admixed with fibrous tissue. It may be monostotic, involving one bone or polyostotic involving several bones.

**Synonyms**
Benign fibro-osseous lesion.

**Epidemiology**
FD affects children and adults and there is no geographical, or racial predilection. The monostotic form affects both sexes equally; the polyostotic form is more common in females by a 3:1 ratio.

**Etiology**
Exact etiology is uncertain. The most recent attempts to define the disorder have focused on genetics and molecular biology.

**Localization**
Any bone in the body can be affected. In the head and neck the skull and facial bones are affected in 10-20% of cases of monostotic disease and 50% of polyostotic cases. In cases with involvement of the temporal bone, the disease is predominantly monostotic. The tympanic, mastoid, squamous or petrous temporal bone may be involved. Other unusual sites include the internal auditory canal, the lateral semi-circular canal and the ossicles. In a retrospective analysis of patients with fibrous dysplasia affecting the skull base, Lustig et al found the temporal bone to be affected in 24%.

**Clinical features**
The main clinical features of disease affecting the temporal bone are: (i) progressive loss of hearing, mostly conductive but which can be sensorineural and profound in some cases, (ii) temporal bone enlargement with progressive bony occlusion of the external auditory meatus, (iii) facial nerve palsy in some patients when the process affects the seventh cranial nerve, (iv) constriction of the ear canal may result in development of an epidermoid cyst lateral to the tympanic membrane likened to cholesteatoma by Megerian et al (1698).

**Macroscopy**
The affected bone is often expanded and the marrow is replaced by firm grey/tan tissue depending on the proportion of bony, fibrous and cartilaginous elements. There may be cyst formation.

**Histopathology**
The lesion consists of irregular trabeculae of woven bone arising abruptly from a bland spindle cell stroma. The trabeculae may be curved and shaped like letters in the Chinese ideogram and are devoid of a rim of osteoblasts. There is no nuclear atypia and mitoses are few. The proportion of fibrous and bony tissue is variable. The lesion may include benign cartilage. Secondary changes include osteoclast giant cells, foamy histiocytes and aneurysmal bone cyst formation.

**Genetics**
Polyostotic fibrous dysplasia (POFD) may occur in the setting of McCune-Albright syndrome, caused by activating mutations in the complex GNAS locus on chromosome 20 (327,527,577).

**Prognosis and predictive factors**
Fibrous dysplasia has rarely been associated with malignant transformation including osteogenic sarcoma, fibrosarcoma and chondrosarcoma, but the temporal bone is not one of the sites where this change has been described.

![Fig. 7.10 Fibrous dysplasia of temporal bone showing irregular bony trabeculae without a rim of osteoblasts.](image1)

![Fig. 7.11 Location of the GNAS1 gene at chromosome 20q13.2-13.3.](image2)
Osteoma and exostosis

Definition
Benign bony enlargement of the deeper portion of the external auditory meatus. There are two distinct forms. Exostosis is more common than osteoma.

ICD-O code
Osteoma 9180/0

Synonyms
Osteochondroma, osteocartilaginous exostosis.

Etiology
Exostosis appears to be related to trauma such as repeated exposure to cold water; in swimmers there appears to be an association with development of exostoses of the tympanic bone [769]. Exostoses have also been observed in individuals who routinely use stethoscopes, e.g., cardiologists [550]. The etiology of osteoma is not clear.

Localization
Osteoma is a very rare lesion, which is a single, unilateral, spherical mass on a distinct pedicle arising in the region of the tympanosquamous or tympanomastoid suture line. It has only occasionally been described outside the external auditory canal and the middle ear, developing in the mastoids, temporal bone internal auditory canal, glenoid fossa eustachian tube, petrous apex and styloid process.

Exostoses are common, broad-based lesions, often bilateral and symmetrical which are usually situated deeper in the ear canal than osteomas. In the bony portion of the normal external auditory meatus there are no adnexal structures and subcutaneous tissue and perios- teum combine to form a thin layer. Therefore the distance between the epidermal surface and underlying bone is small, which may explain the propensity for exostoses of the tympanic bone to develop in those who swim frequently in cold water [2121].

Clinical features
Symptoms are usually those of ear canal obstruction. Osteoma and exostosis are often associated with infection of the external canal on the tympanic membrane side. Surgical removal may be required to enhance drainage as well as to relieve the conducting hearing loss.

Histopathology
The osteoma is a spherical, pedunculated lesion composed of cortical lamellar bone on the outside overlying trabecular bone with intervening marrow spaces. The trabecular bone may show appositional woven bone formation. Normal squamous epithelium of the ear canal is often seen on the surface. The exostosis does not usually show marrow spaces. Both these lesions are distinct from the recently described benign fibro-osseous lesion of the superficial external canal [2121].

Prognosis and predictive factors
These are benign lesions with no potential for malignant transformation.

Fig. 7.12 A Osteoma of deep external canal. From L. Michaels & H. Hellquist (1711). B Exostosis of deep external canal. Note thin epidermal layer on the exostosis above and on the canal skin below and their proximity to the bone. In deeper sections, the exostosis merges gradually with the deep canal bone without pedunculation.

Fig. 7.13 A Exostosis. Coronal CT scan showing broad-based exostosis of deep external canal. B Osteoma. Axial CT scan showing a pedunculated osteoma of one external canal originating from the bone of deep external canal (arrow).
**Angiolymphoid hyperplasia with eosinophilia**

**Definition**
A benign vascular tumour with well formed, but immature, blood vessels, the majority of which are lined by plump, epithelioid (histiocytoid) endothelial cells. Subcutaneous examples are usually associated with a muscular artery. Most cases have a prominent inflammatory component in which eosinophils are a conspicuous feature.

**Synonyms**
Epithelioid haemangioma (ICD-O 9125/0), nodular angioblastic hyperplasia with eosinophilia and lymphofolliculosis, subcutaneous angioblastic lymphoid hyperplasia with eosinophilia and inflammatory angiomatoid nodule.

**Epidemiology**
There is a wide age range with a peak in the third to fifth decades and women are affected more often than men [759, 1945].

**Etiology**
Whether angiolymphoid hyperplasia with eosinophilia is a reactive lesion rather than a neoplasm is still debated. Features cited as supporting a reactive process include a history of trauma (10% of cases), its relationship around a larger vessel showing evidence of damage and the prominent inflammatory component [759, 1945].

**Localization**
The lesion occurs most frequently on the head, particularly the forehead and scalp (often in the distribution of the superficial temporal artery) and in the skin of the ear and the peri-auricular area. Other common sites are the distal parts of the extremities, especially the digits. Other skin surfaces may be involved and occurrences in oral mucous membranes, pharynx and orbit have been reported [759, 1945]. Deep-seated sites are rare, as are an origin in a large vessel.

**Clinical features**
Most patients present with a nodule which has been present for a year or less; sometimes the lesion may have been present for as long as 15 years. In the skin, including that of the ear, the lesions which are often painful or pruritic, appear as dome shaped erythematous or hyperpigmented papules or nodules which may be excoriated and bleed easily. The pre-excision diagnosis is usually that of an angioma or epidermal cyst. There may be several nodules and these can become chronic and coalesce into confluent plaques. There is very little tendency for spontaneous resolution but systemic spread has never been reported. In some patients there is a peripheral blood eosinophilia.

**Macroscopy**
The lesions are usually 0.5-2.0 cm in diameter; they rarely exceed 5.0 cm. Those lesions which contain blood, resemble a haemangioma but in most cases the appearances are rather non-specific. Sub-cutaneous nodules may resemble a lymph node because of circumscription and a peripheral inflammatory reaction./lymphoid reaction.

**Histopathology**
Histologically, there are both vascular...
and inflammatory cellular components. There is a prominent proliferation of small, capillary sized blood vessels. Often there is a vaguely lobular pattern due to clustering of the capillary sized vessels around a medium sized thicker walled vessel. The vessels are lined by plump, epithelioid (histiocytoid) endothelial cells. The vessels look immature and may lack well-defined lumina; sometimes they appear as solid groups of cells. The nuclei of the endothelial cells are large, there is a finely distributed chromatin pattern and often there are central nucleoli. The cytoplasm may appear vacuolated. An inflammatory cell infiltrate with numerous eosinophils, mast cells and lymphocytes is present, though the numbers of eosinophils may vary considerably from case to case. In the peripheral zones of deeper lesions, formation of lymphoid follicles is often present. In deep seated lesions, there is commonly an associated larger blood vessel, usually a muscular artery; and the lining endothelial cells may also appear epithelioid. Dermal lesions are less well circumscribed and demarcated from the surrounding tissue then deeper lesions.

**Immunoprofile**

The epithelioid endothelial cells express CD31 and Factor VIII. CD34 is also expressed but usually only weakly. Immunostaining for actin can be helpful in demonstrating an intact myopericytic layer around the immature vessels. Mast cells are demonstrated by immunostaining with mast cell tryptase, CD117 (C kit) or IgE (since mast cells bear receptors for IgE).

**Prognosis and predictive factors**

While there is no metastatic potential for this tumour, local recurrences following excision occur in up to one third of patients [1945]. The reasons for this are not clear. The recurrences might be the result of incomplete excision, re-growth from a persisting underlying vascular anomaly or merely a reflection of its neoplastic potential. Whatever the reason, follow-up after complete local excision is indicated.

Angiolympoid hyperplasia must be distinguished from Kimura disease [450, 1322] with which it has been confused in

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**Table 7.2 Clinical and histological features of angiolymphoid hyperplasia with eosinophilia (ALH E) and Kimura disease.**

<table>
<thead>
<tr>
<th>Clinical</th>
<th>Angiolymphoid hyperplasia with eosinophilia</th>
<th>Kimura disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex, Age most often</td>
<td>Women, 3rd and 5th decades</td>
<td>Men, young to middle age</td>
</tr>
<tr>
<td>Geographical</td>
<td>Worldwide</td>
<td>Most common in Far East, occasionally in Europe</td>
</tr>
<tr>
<td>Skin/subcutis</td>
<td>Red brown papules</td>
<td>Large disfiguring masses.</td>
</tr>
<tr>
<td>Common sites</td>
<td>Forehead, scalp, ears</td>
<td>Submandibular, parotid, preauricular.</td>
</tr>
<tr>
<td>Regional lymph nodes</td>
<td>Not involved</td>
<td>Often involved</td>
</tr>
<tr>
<td>Eosinophilia</td>
<td>Sometimes</td>
<td>Almost always</td>
</tr>
<tr>
<td>Raised IgE</td>
<td>Never</td>
<td>Always</td>
</tr>
<tr>
<td>Prognosis</td>
<td>Excellent. May recur.</td>
<td>Excellent. Relapses common</td>
</tr>
</tbody>
</table>

**Histopathology**

<table>
<thead>
<tr>
<th></th>
<th>Angiolymphoid hyperplasia with eosinophilia</th>
<th>Kimura disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small vessels</td>
<td>Numerous, immature. May lack lumina or appear as solid groups. May show relation to a larger vessel.</td>
<td>Numerous. Thin walled. Resemble HEVs No association with a large vessel</td>
</tr>
<tr>
<td>Endothelium</td>
<td>Histiocytoid/epithelioid May be vacuolated.</td>
<td>No special features</td>
</tr>
<tr>
<td>Connective tissue</td>
<td>Variable numbers of eosinophils, lymphocytes Numerous mast cells</td>
<td>Oedematous; rich in eosinophils, plasma cells Numerous mast cells</td>
</tr>
<tr>
<td>Lymphoid follicles</td>
<td>Sometimes present</td>
<td>Always present</td>
</tr>
<tr>
<td>Germinal centres</td>
<td>Normal, activated No IgE on FDCs</td>
<td>Polykaryocytes common May contain eosinophils. May be follicle lysis Deposition of IgE on FDCs</td>
</tr>
</tbody>
</table>

HEVs: high endothelial venules
FDCs: follicular dendritic cells
the past since the two diseases were considered to represent the same disease process (2750).

**Kimura disease**
Kimura disease is a chronic inflammatory condition of unknown etiology which presents as large, deep and often disfiguring, subcutaneous masses in the pre-auricular, parotid and submandibular regions. Often, there is enlargement of regional lymph nodes (1390). Occasionally, only lymph nodes are involved. There is a peripheral blood eosinophilia and raised levels of IgE and the histological features are distinctive (1385).

Kimura disease is endemic in the Far East where it affects predominantly young to middle aged men with an age range of 11-80 years. However, the disease also occurs sporadically in Caucasians in the Western World (1069). Histologically, the subcutaneous masses are found to be composed of lymphoid follicles surrounded by oedematous connective tissue rich in eosinophils and containing numerous thin walled blood vessels resembling high endothelial venules (HEVs). Infiltration of the germinal centres with eosinophils and follicle lysis is a frequent finding as is the presence of polykaryocytes. The polykaryocytes (cells with multilobed nuclei resulting from endoreduplication) in Kimura disease are derived from follicular dendritic cells (1069). There is deposition of IgE on the processes of the follicular dendritic cells and there are also numerous mast cells, the latter well shown by immunostaining with antibody to IgE since mast cells bear receptor for IgE (1067). Plasma cells may also be prominent. The disease is self-limiting with an excellent prognosis, though the lesions may recur.

The etiology is unknown. The raised levels of IgE, the IgE deposition in germinal centres and eosinophilia suggest the disease may be atopic in nature; and that the allergic response is results from lymphocyte-mediated interleukin-5 (680).

**Fig. 7.16** Ear. Angiolymphoid hyperplasia with eosinophilia. **A** Capillary sized vessels in an inflammatory cell infiltrate of lymphocytes and eosinophils. **B** High power view of vessels showing their immature appearance, plump histiocytoid endothelial cells and absence of lumina. Note the surrounding eosinophils. **C** Endothelial cell hyperplasia within the central vessel. There is some associated fibrosis and eosinophils are present in the surrounding inflammatory infiltrate. **D** Numerous mast cells, revealed by their receptor to IgE, surround the medium and small sized blood vessels. **E** Factor VIII immunostaining shows arcades of capillary sized vessels in relation to a central muscular artery (this lesion involved the lip). **F** Immunohistochemistry. Membrane expression of CD31 by the histiocytoid endothelial cells.
Idiopathic pseudocystic chondromalacia

Definition
A non-neoplastic swelling of the pinna resulting from localized accumulation of fluid within elastic cartilage.

Synonym
Endochondral pseudocyst

Epidemiology
The lesion occurs mainly in young and middle-aged adults, although it has been reported in children. Minor degrees of this lesion may be present in any damaged ear cartilage.

Etiology
Minor trauma from repeated rubbing of the auricle may play a part. The fluid may exude from undamaged perichondrial vessels that cannot be absorbed by the damaged perichondrial vessels. Small pseudocysts of the elastic cartilage of the pinna may also be seen in the vicinity of inflammatory or neoplastic lesions of that region.

Localization
This condition occurs in any part of the ear cartilage.

Clinical features
The patient complains of painless swelling of a part of the ear cartilage.

Macroscopy
The gross appearance is one of a localized swelling of the auricular cartilage. The cut surface shows a well-defined cavity in the cartilage which is distended with yellowish watery fluid.

Histopathology
Microscopically the cavity shows a lining of degenerated cartilage on one surface; on the other surface the cartilage is normal.

Immunoprofile
There is no expression in the cells lining the cyst-like spaces for CD 31 or cytokeratins indicating that this is an accumulation of fluid in the elastic cartilage rather than an epithelial cyst or a vascular pseudocyst.

Chondrodermatitis nodularis chronica helicis

Definition
A non-neoplastic ulcerating nodule on the helix of the ear, which always involves the underlying cartilage.

Synonym
Winkler disease

Epidemiology
The condition occurs in the third or fourth decades in both sexes.

Etiology
Scleroderma-like changes in the vessels lead to the obstruction of small arteries of the perichondrium which comprise the primary lesions leading to cartilage necrosis. The acute inflammation and epidermal ulceration are secondary to the nearby cartilage necrosis.

Localization
The lesion occurs in the helix of the auricle, less commonly in the antihelix.

Clinical features
A small exquisitely painful ulcerating nodule forms on the auricle, usually in the superior portion of the helix.

Macroscopy
The nodule on the helix is ulcerated in its centre and shows cornified edges. Extruded necrotic cartilage may be seen in the floor of the ulcer.

Histopathology
There is ulceration of the skin of the auricle and complete necrosis of the superficial region of the elastic cartilage of the auricle. A piece of necrotic cartilage infiltrated by neutrophils and bacterial colonies may be present in the floor of the ulcer. The perichondrium of the elastic cartilage shows obstructive thickening of small arteries. Epidermis at the edge of the ulcerated lesion is hyperplastic.

Prognosis and predictive factors
The lesion is usually cured by surgical removal of the painful nodule.
Cholesterol granuloma and cholesteatoma

Cholesterol granuloma

Definition
Cholesterol granuloma is a foreign body giant cell reaction to crystals of cholesterol deposited in the middle ear cleft. It is accompanied by chronic otitis media.

Etiology
Cholesterol granuloma arises from haemorrhage derived from the inflammatory tissue of cholesterol granuloma, the red cell membranes becoming degenerated to cholesterol.

Localization
The main site of cholesterol granuloma is the middle ear cleft. This includes the tympanic cavity and mastoid air cells. Pneumatized air cells at the apex of the temporal bone may also be the site of an expanding destructive lesion of this type.

Clinical features
The tympanomastoid lesions do not, in themselves, produce symptoms. Symptoms of chronic otitis media may be present, however. Cholesterol granulomas of the petrous apex may grow and even invade the cochlea and into the cerebellopontine angle, producing a tumour like mass with hearing loss and life-threatening symptoms.

Macroscopy
Yellow nodules are seen in tympanic cavity and mastoid in this condition. The petrous apex lesions appear cystic, the contents being altered blood.

Histopathology
The yellow tympanomastoid lesions are composed microscopically of cholesterol crystals (dissolved away to leave empty clefts in paraffin-embedded histological sections) surrounded by foreign body type giant cells and other chronic inflammatory cells. Such cholesterol granulomas are almost always found in the midst of haemorrhage in the middle ear mucosa. Hemosiderin is often present within macrophages among the cells surrounding the cholesterol granuloma. The contents of petrous apex cystic lesions are altered blood, and cholesterol clefts with a foreign body giant cell reaction. The wall of such lesions shows granulation tissue with haemosiderin. Remains of low cuboidal (middle ear) epithelium and bone, representing the wall of a pneumatized air cell, may be seen in biopsies of this condition [49,1062].

Cholesteatoma of the middle ear and petrous apex

Cholesteatoma is a misnomer being neither cholesterol containing nor a neoplasm. Cholesteatoma is a cystic or "open" mass of keratin squames with a living "matrix". Although it is not a neoplastic lesion, especially in the middle ear cleft, it may act like one in that it has a propensity to destroy tissue and to recur after excision.

Acquired cholesteatoma of the middle ear

Definition
A cholesteatoma associated with a perforated tympanic membrane is acquired.

Epidemiology
This entity is seen mainly in older children and young adults.

Etiology
It seems likely that the acquired cholesteatoma is derived from entry of external ear canal epidermis into the middle ear. Most cases are associated with severe otitis media in which entry of stratified squamous epithelium from the external ear epidermis through the tympanic membrane occurs. In some cases, it follows blast injury with perforation of the tympanic membrane at the time of the injury [1377]. Acquired cholesteatoma is also known to follow retraction

Fig. 7.19 Ear cholesterol granuloma. A An intact respiratory-type epithelium overlies the cholesterol clefts and foreign-body type giant cells seen in a cholesterol granuloma. B Innumerable histiocytes are seen adjacent to bone with areas of cholesterol cleft formation and inflammatory response.
Cholesterol granuloma and cholesteatomas

Pocket of the tympanic membrane where it is not due to obstruction of the mouth of a retraction pocket, but rather to the deep ingrowth of a band of stratified squamous epithelium from the fundus of the retraction pocket into the middle ear.

Localization
The main site of origin of this lesion is the upper posterior part of the middle ear.

Clinical features
The patient presents with a foul-smelling aural discharge and conductive hearing loss. On examination of the tympanic membrane there is, in most cases, a perforation of the superior or posterosuperior margin.

Macroscopy
The cholesteatoma is seen as a pearly grey structure in the middle ear cavity associated with severe chronic otitis media.

Histopathology
Acquired cholesteatoma is usually "open" rather than "closed" or cystic. The pearly material of the cholesteatoma consists of dead, fully differentiated anucleate keratin squames. This is the corneal layer of the squamous cell epithelium. As in any normal stratified epithelium there are one to three basal layers of cells above which is a prickle (malpighian or spinous) layer composed of five or six rows of cells with intercellular bridges. The deeper layers of the epithelium of the cholesteatoma matrix frequently show evidence of increased proliferation reflected by down-growths into the underlying sub-epidermal connective tissue.

Immunoprofile
The excessive activity has been confirmed by: (a) the strong expression of cytokeratin 16, a marker for hyperproliferative keratinocytes, by cholesteatoma, but its absence in middle ear and external ear epithelium, except in the annulus region of the external tympanic membrane epithelium [278], (b) the strong expression of MIB-1, an antigen related to Ki-67, which also indicates hyperproliferative activity [2494], (c) counts of silver-stained argyrophil nucleolar organizer regions, a technique which likewise displays proliferative activity, shows significantly larger numbers of these structures in the nuclei of acquired cholesteatoma as compared with those of the epidermis of the deep external auditory meatal skin [2496], (d) acquired cholesteatomatous epithelium shows an abnormally high concentration of IL-1, TGF-alpha, EGF-R and 4F2, all being growth factors [2495] indicating greater growth and differentiating activity than is present in normal epidermis.

Genetics
Acquired cholesteatoma does not show DNA aneuploidy nor does it possess an inherent genetic instability, a critical feature of all malignant lesions [33].

Congenital cholesteatoma of the middle ear

Definition
Congenital cholesteatoma is defined in clinical practice as a cholesteatoma of the middle ear which exists in the presence of an intact tympanic membrane, the implication being that severe chronic otitis media, which normally produces a perforation of the tympanic membrane, has not led to its development.

Epidemiology
This lesion is found in infants and young children.

Etiology
Small colonies of cells confirmed by immunohistochemistry as being epidermoid in nature are found near the tympanic membrane on the lateral anterior superior surface of the middle ear in every temporal bone after 15 weeks gestation. These “epidermoid formations”, are derived from the actively growing epidermis of the eardrum. They increase significantly in size with increasing age and at the same time show increasing epidermoid differentiation [1502]. In normal development, the epidermoid colonies disappear by the first post partum year. However, if one of them does not resolve, but continues to grow, this will become a congenital cholesteatoma.

Localization
The majority of cases are found in the antero-superior part of the middle ear.

Clinical features
In early lesions there are no symptoms,
the cholesteatoma being discovered by routine otoscopy. In later cases, the lesion is much larger and symptoms may resemble those of acquired cholesteatoma.

**Macroscopy**
In most cases a spherical whitish cyst in the anterosuperior part of the tympanic cavity is seen, behind an intact tympanic membrane. In 10% of congenital cholesteatomas the lesion is “open”, the desquamated squames entering the tympanic cavity.

**Histopathology**
The matrix of congenital cholesteatoma is epidermis, comprising a single row of basal cells, several rows of malpighian cells and a thin granular layer. The surface of dead, keratinous squames merges with the keratinous contents of the cyst, or keratinous lamellae in the case of the open type.

**Immunoprofile**
Immunostaining shows similar features to those of acquired cholesteatoma.

**Prognosis and predictive factors**
If removed early, when small, congenital cholesteatoma can be considered as cured. If left, or not diagnosed, until later in life, the problems of middle ear damage and recurrence become similar to those of acquired cholesteatoma.

**Cholesteatoma of the petrous apex**

**Definition**
An epidermoid cyst arising in the region of the petrous apex. It bears no relation to cholesteatoma of the middle ear.

**Etiology**
It is probably of congenital origin, but no cell rest has been discovered from which it might arise.

**Clinical features**
This lesion usually presents with facial palsy and hearing loss, due to involvement of the seventh and eighth cranial nerves, respectively, in the cerebellopontine angle (564).

**Histopathology**
The histological appearance is similar to that of middle ear cholesteatomas.
Adenoma of the middle ear

Definition
Adenoma is a benign glandular neoplasm showing variable differentiation along neuroendocrine and mucin-secreting pathways.

ICD-O code 8140/0

Synonyms
Middle ear adenomatous tumour, neuroendocrine adenoma of the middle ear, carcinoid of the middle ear.

Epidemiology
This is an uncommon neoplasm, but among the most frequent ones arising in the middle ear. There is an approximately equal sex distribution, with an age range of 20-80 years, and a mean age of 45 years [2623].

Localization
The tumour arises anywhere in the middle ear cavity, sometimes extending into the mastoid. In one reported case it arose from the epitympanic part of the tympanic membrane [75]. In a small number of cases it may be found to have spread through the tympanic membrane (2623).

Clinical features
Patients complain of muffled hearing with a pressure sensation in the affected ear. Otoscopy shows an intact tympanic membrane in the first stage with a dark brown-reddish coloured structure behind it. Tumour may later expand and involve the ossicular chain causing conductive hearing loss and may penetrate the tympanic membrane. Treatment is surgical. The tumour is usually easily removed, but if ossicles are entrapped reconstructive surgery is needed.

Macroscopy
The neoplasm has been described as being white, yellow, grey or reddish brown at operation and, unlike paraganglioma, is usually not vascular. Although not encapsulated it seems to peel away from the walls of the surrounding middle ear with ease, although ossicles may sometimes be surrounded by the tumour and may even show destruction.

Histopathology
Adenoma is formed by closely apposed small glands with a “back to back” appearance. In some places a solid or trabecular arrangement is present. Sheet-like, disorganized areas are seen in which the glandular pattern appears to be lost. This may be artefactual and related to the effects of the trauma used in taking the biopsy specimen, on the delicate structure of the cells, but the appearance may erroneously lead one to suspect malignancy. The cells are regular, cuboidal or columnar and may enclose luminal secretion. A distinct and predominant "plasmacytoid" appearance of the epithelial cells of the neoplasm may be displayed [2164]. The small central nuclei rarely contain nucleoli and show no significant mitotic activity. No myoepithelial layer is seen. Periodic acid-Schiff and Alcian blue stains may be positive for mucoprotein secretion in the gland lumina and in the cytoplasm of the tumour cells. Soon after adenoma of the middle ear was described in 1976 [588,1160], it was reported that some glandular tumours of the middle ear, otherwise apparently identical to an adenoma, showed neuroendocrine features as shown by Grimelius positivity, the presence of numerous membrane-bound granules on electron microscopy, and expression of immunohistochemical markers for neuroendocrine activity. The concept of “carcinoid tumour” evolved, i.e. that this was a distinct neoplasm with significant neuroendocrine differentiation. As with carcinoids in other locations, it was considered to have malignant potential. It is now clear that most, probably all, middle ear adenomas express neuroendocrine markers (2623,2727).

Immunoprofile
Neuroendocrine markers such as synaptophysin, chromogranin, and various polypeptides, are demonstrated in addition to cytokeratins [2623].

Electron microscopy
Ultrastructural examination of five cases showed basally situated cells and solid tumour containing neuroendocrine granules which were positive for neuroendocrine markers. This is in contrast to apically situated dark cells which contained mucous granules and were negative for neuroendocrine markers [2727].

Precursor lesions
The lesions arise from the lining epithelium of the middle ear. Under appropriate stimuli such as otitis media, this epithelium has the potential for glandular differentiation. However, neuroendocrine differentiation has not been demonstrated in either normal or “metaplastic” glandular epithelium.

Genetics
There has so far been no study of molecular genetic aspects of this tumour. This neoplasm does not occur in families.

Prognosis and predictive factors
There have been a few recurrences after incomplete local surgical excision.
Papillary tumours of the middle ear

L. Michaels
A. Sandison
G.L. Davis

Aggressive papillary tumour

Definitions
Tumour with a papillary, non-stratified epithelial pattern showing invasive behaviour.

ICD-O code 8260/1

Synonyms
Primary adenocarcinoma of the middle ear of papillary type, aggressive papillary tumour of temporal bone, papillary adenoma.

Epidemiology
Forty-six cases with this neoplasm were collected from the literature in 1994 (843). Some of these had been reported as low-grade adenocarcinoma of probable endolymphatic sac origin (1038). Review of each of the case reports in these two studies, together with cases reported more recently, reveals a total of 24 cases in which the middle ear was involved, comprising 17 females and 7 males. The age-range at time of diagnosis was between 16 and 55 years with a median age of 33 and a mean age of 34 years. In many of the cases, however, the patient had already suffered symptoms subsequently ascribable to the tumour for some years when the diagnosis was made, so that the age of onset may be considerably younger than is suggested.

Localization
The tumour is found in any area of the middle ear, including the mastoid process and air cells and may fill the tympanic cavity. In all of the described cases, except two (519,2481) there was extensive invasion outside the middle ear, involving the apical portion of the petrous bone in most and in a few the tumour reached the cerebellopontine angle and the cerebellum. It has been suggested that cases of aggressive papillary middle ear tumour with widespread involvement of the temporal bone may arise from a primary papillary adenocarcinoma of the endolymphatic sac (1038). The frequent association of papillary tumours in the middle ear with apical petrous bone neoplasia of the same type, the similarity of the histological appearances of the neoplasms in the two regions and the association of some cases of papillary tumours in both regions with von Hippel-Lindau disease would seem to favour this concept, but an origin in the middle ear in some cases of this neoplasm has not been definitely excluded. This would explain the presence of the neoplasm in the middle ear only in two described cases. In the single description of the pathological changes of aggressive papillary tumour of the middle ear in an autopsied temporal bone, widespread deposits of tumour at inner ear sites are depicted, but no mention is made of involvement of the endolymphatic sac or duct (2355). Whatever the site or sites of origin of this tumour it should be recognized that papillary epithelial tumour of the middle ear is frequently an aggressive neoplasm, in contrast to the non-papillary adenoma of the middle ear which is quite benign (1741).

Clinical features
In most cases of this neoplasm clinical and audiological features point to a middle ear lesion. Suspicion of a neoplasm of the middle ear is enhanced by the otoscopic features in a few cases. Indeed, the tympanic membrane has been perforated by the tumour, which is seen to lie in the external canal in some cases. On imaging the medial parts of the petrous temporal bone show, in the great majority of cases, areas of involvement by a lytic lesion, representing an invasive neoplasm which may extend posteriorly outside the temporal bone and invade the cerebellum (843). Fifteen percent of patients with aggressive papillary tumour of middle ear have been found to possess neoplasms or other manifestations of Von Hippel-Lindau syndrome. There may be a family history of this condition in the patient without its actual physical manifestations (843).

Histopathology
The middle ear cleft, including the mastoid air cells, is usually filled with the papillary tumour. Bone invasion is often seen. A papillary glandular pattern is present...
Papillary tumours of the middle ear

with complex interdigitating papillae lying loosely or infiltrating fibrous connective tissue. The papillae are lined by a single layer of low, cuboidal to columnar epithelial cells with uniform nuclei, eosinophilic cytoplasm and indistinct cell borders. Thyroid follicle-like areas may be present, similar to those seen in endolymphatic sac carcinoma.

**Immunoprofile**
Markers for cytokeratin, epithelial membrane antigen and S100 are positive. The absence of thyroglobulin must be determined to exclude metastatic papillary carcinoma of the thyroid. Markers for CK7, CK20 and carcinoembryonic antigen may also be useful to exclude metastatic deposits from lung and colon.

**Genetics**
The genetic aspects of von Hippel-Lindau disease are described below. In view of the association of some cases of that condition with aggressive papillary middle ear tumours it is suggested that the clinical assessment of each case with the latter neoplasm should include an investigation for the gene mutations of von Hippel-Lindau disease.

**Schneiderian-type papilloma**
Schneiderian epithelium refers to the normal respiratory-type ciliated epithelium of the nose and paranasal sinuses. Schneiderian papillomas are tumours of the nose and paranasal sinuses that are stated to be derived from this epithelium. Three such types of papillomas are described: inverted (endophytic), exophytic (fungiform, everted) and oncocytic (cylindric cell). Intermediate types are said to be found between the three forms (1741). It has, however, been denied that such intermediate forms exist and it is suggested that the three types are each separate and distinct entities of the nose and paranasal sinuses (1714). Of the three histological forms only inverted papilloma is characteristically a sinonasal neoplasm. The other two types of Schneiderian-type papilloma may be seen at other sites. Low-grade squamous carcinoma in the nose may sometimes be mistaken for inverted papilloma (1710).

Fourteen cases of middle ear tumours purportedly resembling Schneiderian-type papilloma have been found in the literature. Each of these cases is listed in Table 7.3; wherever possible the histological appearances are summarized in the table; in some insufficient or no histological description was given. In two cases only were the features of inverted papilloma depicted: in Case 1 the term “inverted” and in Case 2 the term “endophytic” were used to describe the neoplasm. “Inverted” or “endophytic” features comprised only a portion of the tumours in the two cases. In Case 2 (1242) the term transitional cell papilloma was used. This is a term that has frequently been applied to describe everted squamous cell papilloma. In this case inverted papilloma was found in the nasal cavity and it was suggested that the papillomas might have spread from there to the middle ear by way of the

<table>
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<td>“Inverted papilloma” and high grade carcinoma</td>
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<td>2. Kaddour et al (1242)</td>
<td>“Transitional cell papilloma” Inverted papilloma in nose</td>
<td>Inverted papilloma derived from nasal tumour</td>
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<td>3. Roberts et al (2184)</td>
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<td>4. Seshul et al (2308)</td>
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<td>5. Wenig (2757) Case 1</td>
<td>Epidermoid papilloma” with “inverted” and “cylindric cell papilloma”</td>
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<td>6. Wenig (2757) Case 2</td>
<td>“Epidermoid papilloma” with exophytic and endophytic growth</td>
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<td>7. Wenig (2757) Case 3</td>
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<td>8. Wenig (2757) Case 4</td>
<td>“Epidermoid papilloma” with features of “cylindric cell papilloma”</td>
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<tr>
<td>9. Wenig (2757) Case 5</td>
<td>“Epidermoid papilloma” with features of “cylindric cell papilloma”</td>
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<tr>
<td>10. Jones et al (1234)</td>
<td>Squamous epithelium with areas of carcinoma in-situ</td>
<td>Squamous carcinoma of middle ear</td>
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<td>11. Chhetri et al (423)</td>
<td>Papillary with areas of squamous epithelium adjacent to respiratory epithelium</td>
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</table>
Eustachian tube. In Cases 1, 4 and 10 “inverted papilloma” were found in the middle ear concomitantly with in situ or invasive squamous carcinoma and it seems possible that the “inverted papilloma” areas might have been, in reality, areas of low grade squamous carcinoma.

We would suggest that a good case has not been made for the occurrence of inverted papilloma in the middle ear. Some of the lesions may have been papillomas of the middle ear as described above. In Case 2 inverted papilloma could conceivably have colonized the middle ear from the nasal cavity. Further detailed descriptions of the entity are required to justify the diagnosis of such a diagnostic category in this situation.

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**Inverted papilloma**

**Definition**

Papillary neoplasm of the middle ear which is histologically identical to that occurring in the sinonasal region.

**ICD-O code** 8121/1

**Localization**

Middle ear. Also occurs in association with similar papillomas of the upper respiratory tract, either by direct continuity or in a multicentric fashion.

**Clinical features**

It usually presents as chronic otitis media.

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**Macroscopy**

Polypoid tumour filling the middle ear cavity.

**Prognosis and predictive factors**

Thus far, these tumours have shown no evidence of invasion but recurrences are common [2757].

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**Choristoma**

**Definition**

A choristoma in contrast to a hamartoma, is composed of tissues which are not normally present in the part of the body where it is found. Choristomas are occasionally seen in the middle ear. They are composed of one or other of two types of tissue: salivary gland or glial tissue.

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**Salivary gland choristoma**

**Localization**

This lesion usually occurs as a mass in the middle ear attached posteriorly in the region of the oval window. There are usually absent or malformed ossicles [1093].

**Histopathology**

Salivary gland choristomas consist as a rule of lobulated mixed mucous and serous elements like the normal submandibular or sublingual gland, but unlike the parotid gland.

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**Glial choristoma**

**Clinical features**

In this lesion, masses composed of glial tissue are identified in biopsy material from the middle ear. A bony deficit with consequent herniation of brain tissue into the middle ear should be ruled out by imaging [1257].

**Histopathology**

Glial masses are present, composed largely of astrocytic cells with large amounts of glial fibrils.

**Immunoprofile**

The identity of the tissue as glial may be confirmed by immunohistochemical staining for glial acidic fibrillary protein.
Squamous cell carcinoma of the middle ear

**Definition**
A malignant tumour composed of stratified squamous epithelium arising from the cuboidal and / or pseudostratified epithelium of the middle ear.

**ICD-O code**
8070/3

**Synonyms**
Epidermoid carcinoma, squamous carcinoma

**Epidemiology**
The disease affects males and females equally, with an age range of 34-85 years and an average age of 60 years.

**Etiology**
An origin from long-term chronic inflammation of the middle ear has been suggested. However, malignant neoplasia in its earlier stages has clinical features similar to those of chronic otitis media. Moreover biopsy is not usually carried out during surgery when a diagnosis only of otitis media has been made. Therefore, longstanding squamous carcinoma of the middle ear may go undiagnosed.

**Localization**
The neoplasm soon expands to involve much of the middle ear. There is extension by tumour through the bone on the medial wall of the Eustachian tube to infiltrate the perineurium of nerves in the carotid canal. The tumour also penetrates the thin layer of bone between the posterior mastoid air cells and the dura with subsequent invasion along the dura and into the internal auditory meatus. Bilateral squamous cell carcinomas of the middle ear have been described [1713].

**Clinical features**
This tumour is usually advanced at presentation. The patient usually complains of pain in the ear, bleeding and a serosanguinous discharge from the ear canal. In those cases with a concomitant external canal carcinoma a plaque-like or even polypoid mass may be felt or even seen in the canal. Seventh nerve palsy is an important sign indicating infiltration beyond the middle ear.

**Macroscopy**
A tumour fills the middle ear and may extend into the mastoid air cells and along the pathways described above.

**Histopathology**
The neoplasm is a keratinizing squamous cell carcinoma with a variable degree of differentiation. Atypical change and even carcinoma in situ may be seen in some parts of the middle ear epithelium adjacent to the tumour. The tumour arises from malignant stratified squamous epithelium and in certain areas an origin directly from basal layers of cuboidal or columnar epithelium may be seen.

**Precursor lesions**
There is no evidence of a relationship to cholesteatoma or the epidermoid cell rests which normally occur in the middle ear during development.

**Prognosis and predictive factors**
The prognosis is uniformly poor and does not correlate with degree of tumour differentiation.

---

Fig. 7.26 Squamous carcinoma. Autopsy temporal bone specimen showing infiltration of squamous carcinoma of middle ear into mastoid air cell.
Meningioma of the middle ear

L. Michaels
S. Soucek

Definition
Meningioma is a benign tumour usually forming intracerebrally, but sometimes seen involving bony structures around the brain including the middle ear. It arises from the pia-arachnoid cells of the meninges.

ICD-O code
9530/0

Epidemiology
Meningioma of the middle ear affects women more than men, shows an age range of between 10 and 80 years with a mean age of 49.6 years. Female patients present at an older age (women 52.0 years, men 44.8 years) [2597].

Localization
Meningiomas occur at a number of sites in the temporal bone, including the internal auditory meatus, the jugular foramen, the geniculate ganglion region and the roof of the Eustachian tube [1830]. The most common temporal bone site for primary meningioma is in the middle ear cleft. In a recent study (36 patients), most tumours involved the middle ear, but a few involved adjacent structures such as the external canal or temporal bone. Only two showed extension from CNS on imaging [2597].

In neurofibromatosis type 2 (NF-2 see below), meningioma-like masses occur commonly in the internal auditory canal and cerebellopontine angle.

Clinical features
Patients present clinically with hearing change, otitis media, pain, and/or dizziness / vertigo.

Macroscopy
Gross appearances are those of a granular mass with a gritty consistency.

Histopathology
Microscopically the neoplasm in the middle ear shows the same histological features of any of the well-described subtypes of intracranial meningioma. The most common variety seen in the middle ear is the meningothelial type, in which the tumour cells form masses of epithelioid, regular cells often disposed into whorls. Occasionally, fibroblastic and psammomatous variants are seen.

Immunoprofile
Histological diagnosis may be difficult because the typical features of meningioma are absent. Under these circumstances immunocytochemistry is of diagnostic value. Vimentin and epithelial membrane antigen are expressed in the majority of meningiomas and cytokeratins are uniformly negative. Expression of S-100 protein identifies spindle cell tumours as of neurogenic origin, thus excluding spindle cell meningioma.

Prognosis and predictive factors
Although Nager's review of temporal bone meningiomas (1964) indicated that only two out of 30 patients survived a 5-year period [1830], more recent experience of middle ear meningiomas signals a better outlook after careful local excision (5-y survival, 83%).

Fig. 7.27 Meningioma of middle ear. A Meningioma of middle ear, meningothelial type, showing small whorls. B Ear meningioma. A variety of different growth patterns can be seen, but the meningothelial nature of the neoplasm is always maintained.
Vestibular schwannoma

Definition
A benign nerve sheath tumour arising in the internal auditory canal.

ICD-O code 9560/0

Synonyms
Acoustic neuroma, acoustic neurinoma, neurilemmoma

Epidemiology
Vestibular schwannoma is the most common neoplasm of the temporal bone. Unilateral vestibular schwannoma accounts for 5-10% of all intracranial tumours and for most of the cerebello-pontine angle tumours. It is found in about 0.8% of consecutive adult necropsies (1475). The age at presentation is the fifth or sixth decade. It also is seen in younger people in association with neurofibromatosis type 2.

Etiology
Solitary vestibular schwannoma occurs sporadically, and does not seem to be associated with a gene mutation. The etiology is unknown.

Localization
Vestibular schwannoma was formerly considered to arise most commonly at the glial-neurilemmal junction of the eighth cranial nerve. Such a site of origin has now become doubtful (2834). In one study of five temporal bones with small vestibular schwannomas, the tumour arose more peripherally (2834). The vestibular division of the nerve is usually affected. Rarely, the cochlear division is the source of the neoplasm. Growth takes place from the site of origin of the tumour, both centrally onto the cerebello-pontine angle and peripherally along the canal. Vestibular schwannoma is usually unilateral, but may be bilateral, in which case the condition is neurofibromatosis 2.

Clinical features
Progressive unilateral hearing loss (90% of patients) and tinnitus (70% of patients) are the clinical manifestations, due to cochlear involvement. Less common symptoms are: headache, vertigo, facial pain and facial weakness. The neoplasm may grow slowly for years without causing symptoms and may be first diagnosed only at post-mortem. Diagnosis is usually made by MRI scanning. In small, slowly growing tumours, an option for management is non-surgical, using MRI scanning at intervals to observe growth. Surgical removal may be carried out by drilling from the external canal through the temporal bone or by craniotomy and middle fossa approach to the internal auditory meatus, or by stereotactically guided gamma knife surgery.

Macroscopy
The neoplasm is of variable size and shape. Small tumours either do not widen the canal at all or produce only a small indentation in the bone.

Fig. 7.28 Vestibular schwannoma. A Microsliced autopsy temporal bone. The neoplasm arises from the vestibular division of the eighth nerve and compresses the cochlear division. Note the granular deposit lining the cochlea, a feature of most larger vestibular schwannomas. From L. Michaels & H. Hellquist (1711). B Axial T2 post-contrast MRI scan of the posterior fossa showing a well-defined intracanalicular vestibular schwannoma (arrow). Note the eighth cranial nerve leading into the tumour.
**Immunoprofile**
These tumours express diffuse, strong nuclear positivity for S-100 protein. Vimentin expression is also usually positive though not specific. These findings are common to both unilateral vestibular schwannoma and the schwannomas of NF2. Glial fibrillary acidic protein and neuron specific enolase markers may also be expressed. The tumours are consistently negative for CD34, a marker widely used for the diagnosis of solitary fibrous tumours, unless there are widespread degenerative changes [2625]. The proliferation marker Ki67 has demonstrated that tumours 18 mm or smaller in diameter have lower proliferation indices and growth rates than tumours larger than 18 mm [186].

**Genetics**
Ninety five per cent of vestibular schwannomas are unilateral and are sporadic. Less than 5% of tumours are bilateral and therefore associated with the NF2 gene. The risk that a unilateral tumour is the first indication of NF2 is closely related to the age of the patient. Young patients under the age of 25 are at high risk of developing contralateral tumours and NF2 while patients with unilateral tumours who are over the age of 55 have virtually no risk of developing NF2. There is no increased incidence of unilateral vestibular schwannoma or NF2 in the offspring of patients with unilateral vestibular schwannoma [1595].

**Electron microscopy**
Schwann cells are identified as the cell of origin by their Interdigitating slender cytoplasmic processes covered with a continuous layer of basal lamina [434]. Extensive degeneration of the vestibular sensory organ as detected ultrastructurally is brought about by growth of the neoplasm from the vestibular division of the eighth nerve. Even small tumours may cause this change [2241].

**Microneuromas and Paget disease of bone**
**Definition**
Small non-neoplastic tumours composed of masses of intertwined bundles of nerve fibres and Schwann cells, which are sometimes observed near the cochlea or vestibule in the temporal bones of cases of Paget disease of bone. It is likely that they are the result of pressure by the growth of the pagetic bone on the nerve fibres with their regeneration and the production of traumatic neuromas [2275].

**Prognosis and predictive factors**
Size is an important aspect in the prognosis of cases of vestibular schwannoma. Tumours 18 mm or smaller in diameter have lower proliferation indices and growth rates than tumours larger than 18 mm [186].

---

**Fig. 7.29 Vestibular schwannoma.**

A Small vestibular schwannoma (S) in autopsy temporal bone specimen. It is arising from the vestibular division of the eighth nerve and causing a small indentation only of the bony wall of the internal canal. There is exudate in the vestibule, but not in the cochlea. From L. Michaels & H. Hellquist (1711). B Vestibular schwannoma showing Antoni A appearance and hyaline blood vessels.
Neurofibromatosis type 2

**Definition**
An autosomal dominant disorder characterized by a high incidence of bilateral vestibular schwannomas as well as schwannomas of other cranial and peripheral nerves, and other benign intracranial and intraspinal tumours.

**Synonym**
Bilateral acoustic neuroma, bilateral vestibular schwannoma

**Epidemiology**
The condition usually presents clinically in the first or second decade of life.

**Localization**
Bilateral vestibular schwannoma is characteristic of neurofibromatosis type 2. The tumours usually arise from the superior vestibular branch of the 8th cranial nerve. In addition, schwannomas of other cranial and peripheral nerves do occur as well as a wide variety of other benign intracranial and intraspinal tumours including schwannoma of other cranial and peripheral nerves, meningiomas, ependymomas, spinal neurofibromas, and gliomas. Juvenile posterior subcapsular cataract is also found.

**Clinical features**
Vestibular schwannomas in NF2 patients grow more rapidly than sporadic unilateral tumours. Infiltration of the cochlear and facial nerves occurs, making it more difficult to preserve hearing and facial nerve function after surgery. Screening of the relatives of affected subjects is necessary. Affected relatives of these patients often have normal audiograms and normal auditory brain stem responses in the presence of a schwanna, and it has been recommended that the screening of relatives of NF2 patients should be by magnetic resonance image scanning with gadolinium (Gd-DTPA) enhancement [1326]. In contrast to NF2, neurofibromatosis type 1 or von Recklinghausen disease is characterized by dermal neurofibromas. However, otological manifestations of neurofibromatosis were recorded in 6.5% of children with NF1, involving external ear canal, middle ear and eighth cranial nerve [2415].

**Macroscopy**
The gross appearance of the vestibular schwannomas in neurofibromatosis 2 is similar to that of sporadic vestibular schwannoma.

**Tumour spread and staging**
There is invasion of the facial nerve in the internal auditory canal and also invasion of the modiolus and bony vestibular wall in some cases [2354].

**Histopathology**
Vestibular schwannomas in NF2 are similar to those of sporadic vestibular schwannoma.

**Immunoprofile**
The degree of labelling with the proliferation marker Ki67 is higher in cases of NF2 that in those of solitary vestibular schwannoma [17]. Otherwise, the immunoprofile is identical.

**Genetics**
NF2 is an autosomal dominant condition. About 50% of patients have NF2 as a result of a new mutation and 50% inherit the disease from an affected parent. The children of an affected person have a 50% chance of inheriting the disease and prenatal diagnosis is available. The gene for NF2 is a suppressor gene which has been mapped to the long arm of chromosome 22 (22q12). It codes for a protein which has been called by two names; MERLIN which stands for moeizin-ezrin radixin like protein because it resembles the family of cytoskeletal associated proteins and SCHWAN-NOMIN because of its role in preventing schwannoma formation. It is a membrane-associated protein believed to inhibit cell growth and motility and preserve cell shape as well as anchoring the cell cytoskeleton to the surrounding matrix. Studies aimed at identifying germline mutations in patients with NF2 found mutations in up to 2/3 of cases. A wide variety of mutations have been found in all exons of the NF2 gene apart from exons 16 and 17. Ninety per cent of the mutations are predicted to truncate the gene product by introducing a stop codon, a frameshift with premature termination or a splicing alteration. This suggests that loss of the protein’s function is necessary for tumourigenesis [1595]. A family on the Isle of Man, Great Britain, with inherited salivary gland neuroendocrine carcinoma and amelogenesis imperfecta [1712] also displayed vestibular schwannoma in two male sibs, bilateral in one. Molecular genetic analysis has not yet been carried out, but it is likely that the disease process in this family is genetically related to NF2.
Lipoma of the internal auditory canal

Definition
A benign tumour of adipocytes, important in this situation because it can mimic vestibular schwannoma.

ICD-O code 8850/0

Localization
Either in the cerebellopontine angle or in the internal auditory canal.

Clinical features
The most frequent associated symptoms are of cochleovestibular origin, such as hearing loss, dizziness and unilateral tinnitus. Other associated symptoms involve the facial nerve or the trigeminal nerve. Complete resection is associated with cranial nerve damage. The lesion may be mistaken clinically for a schwannoma, but magnetic resonance imaging can distinguish between the two entities. Where there is doubt, frozen section diagnosis of an incisional biopsy should be carried out to avoid the risk of damage to the 7th or 8th cranial nerve or their branches which may pass through the tumour.

Macroscopy
There may be erosion of the walls of the internal auditory canal as with vestibular schwannoma, and lipoma may appear similar to the latter at operation.

Histopathology
The tumour is similar to lipomas elsewhere except that 7th or 8th cranial nerve or their branches may be present among the adipocytes.

Haemangioma

Definition
A benign tumour of blood vessels.

ICD-O code 9120/0

Synonyms
Cavernous haemangioma, vascular tumour, vascular malformation.

Etiology
Little is known about these rare tumours. Only 43 cases were reported in the world literature up to the year 2000. They are thought to arise from the dense vascular networks around the geniculate ganglion and Scarpa ganglion, which may account for the site predilection.

Localization
Haemangioma of the temporal bone occur most frequently at two sites, the internal auditory meatus and the geniculate ganglion. Cavernous haemangioma arising in the cerebellopontine angle can mimic vestibular schwannoma, which occur more commonly at this site.

Clinical features
Patients may present with hearing loss or facial paralysis due to VIIth cranial nerve involvement, which usually happens at an early stage, before the tumour reaches 1 cm diameter. Symptoms are suggestive of haemangioma if they are disproportionate to the size of the lesion as seen on imaging, or fluctuate with hormonal changes such as occur in pregnancy. MRI imaging shows a lesion that enhances with gadolinium and which may contain areas of microcalcification.

Histopathology
The lesions are composed of irregular dilated vascular spaces with collagenous walls lined by a single layer of endothelium.

Prognosis and predictive factors
Although they are benign lesions, haemangiomas enlarge and do not spontaneously involute. Early surgical intervention is recommended to best preserve facial nerve function.
**Endolymphatic sac tumour**

**Definition**
Endolymphatic sac tumour (ELST), a non-metastasizing adenocarcinoma of endolymphatic sac origin, is a slowly-growing tumour which widely invades the petrous bone.

**ICD-O code**
8140/3

**Synonyms**
Heffner tumour (1038), low grade adenocarcinoma of endolymphatic sac origin (LGAES), aggressive papillary middle ear tumour (APMET).

**Epidemiology**
A rare neoplasm of adults. Although ELST is extremely rare in the general population. An association with von Hippel-Lindau disease (VHL) is established (1696).

**Etiology**
The mutations in the VHL gene have been implicated in the development of ELST.

**Localization**
At an early stage of its growth, the neoplasm is sited within the endolymphatic sac (970,1026). At a later stage, it destroys much of the petrous bone, including the middle ear and extends to the posterior fossa into the cerebellopontine angle (1038,2767).

**Clinical features**
Tinnitus, hearing loss and vertigo, similar or identical to the symptoms of Ménière disease, are present in about one third of patients. It is presumed that early obstruction of the endolymphatic sac leads to hydrops of the endolymphatic system of the labyrinth. As the tumour spreads, facial nerve paralysis and/or cerebellar disorders may develop. Imaging reveals a lytic temporal bone lesion, appearing to originate from the region between the internal auditory canal and sigmoid sinus (which is the approximate position of the endolymphatic sac). There is eventually prominent extension into the posterior cranial cavity and invasion of the middle ear.

**Histopathology**
In most cases, ELST has a variable papillary-glandular appearance, the papillary proliferation being covered by a single row of low cuboidal cells. The vascular nature of the papillae in some cases has given the tumour a histological resemblance to choroid plexus papilloma. Some cases show areas of dilated glands containing secretion which resembles colloid. Such thyroid-like areas may even dominate the histological pattern. A few cases show a clear cell predominance resembling carcinoma of the prostate and renal cell carcinoma. Despite controversy surrounding the origin of so-called “aggressive papillary middle ear tumour” (APMET see above) (844), current evidence suggests that it is ELST with extension into the middle ear.

**Immunoprofile**
These tumours express cytokeratin and some express glial fibrillary acidic protein. Specific markers for metastases including thyroglobulin and prostate-specific antigen are negative.

**Genetics**
The gene responsible for Von Hippel-Lindau (VHL) has been mapped to the short arm of chromosome 3 (3p 25-26). Mutations in this gene have been reported in patients with ELST. The gene is thought to be a tumour suppressor gene since genetic analysis of tumours in patients with VHL disease supported Knudson’s hypothesis that an inherited mutation in one allele, followed by somatic mutation and loss of function of the second allele were required for tumorigenesis. In the case of sporadic tumours, tumourigenesis results from somatic mutation in both alleles of the tumour suppressor gene. A germline mutation of the VHL gene and somatic mutation of the wild type allele have been shown in ELST from VHL patients and somatic mutations in the VHL gene have been demonstrated in sporadic ELST. The rarity of ELST has meant that analysis of the tumours for specific mutations has been difficult. Hamazaki et al reported the genetic analysis of a case of ELST which

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**Fig. 7.34** Endolymphatic sac tumour. Axial T2 weighted MRI scan of the posterior fossa. The white area represents fluid within the endolymphatic sac and the grey area the solid component of the endolymphatic sac tumour.

**Fig. 7.35** Endolymphatic sac tumour. **A** Mild atypical nuclei are identified at the luminal surface of the papillary projections. Thin fibrovascular cores are present. **B** Endolymphatic sac tumour showing thyroid-like glandular pattern.
occurred in the absence of VHL disease (984). This showed a nucleic acid substitution of G to T in nucleotide 546 in the VHL gene which resulted in an amino acid substitution (Trp to Cys codon 117). An identical mutation has been reported in other VHL families. This suggests that the VHL gene is associated with ELST tumourigenesis with or without VHL disease.

**Prognosis and predictive factors**

The tumour grows slowly over many years and is not known to metastasise. Many tumours have already attained a large size at presentation. It is important to screen all patients with VHL for ELST by imaging so that small tumours may be detected early and completely excised (1696). Likewise, all patients with ELST should be screened for the VHL gene.
These tumours, in the WHO classification of haematological malignancies (1197), are primarily stratified according to lineage into myeloid, lymphoid, histiocytic/dendritic cell and mast cell neoplasms. In each category, neoplasms are defined according to morphology, immunophenotype, genetic features and clinical syndromes. For each type of neoplasm a cell of origin is proposed.

**Ear lymphomas**
Lymphomas occurring in and around the ear are rare compared to other sites. They may involve the pre- and retro-auricular lymph nodes, temporal bone or skin and soft tissue. Those lymphomas affecting the pre-auricular lymph nodes are predominantly disseminated or nodal. Lymphomas of bone tend to occur with persistence of red marrow. With the exception of plasma cell tumours such as plasma cell myeloma (synonyms: multiple myeloma, myelomatosis) and plasmacytoma (synonym: solitary plasmacytoma of bone), both of which can involve the squamous and petrous temporal bone, lymphomas are extremely rare in the temporal bone. The mastoid process, part of the temporal bone, contains air cells and lacks marrow.

**B-cell chronic lymphocytic leukaemia/small lymphocytic lymphoma (B-CLL / SLL)**

**ICD-O code** 9823/3

Of those lymphomas resulting in cutaneous lesions of the head and neck, including the ears, the most common is B-cell chronic lymphocytic leukaemia/small lymphocytic lymphoma. Leukaemic infiltrates in the skin or leukaemia cutis are not rare. Lesions on the face including ears have been recognized for several decades as heralding the onset of B-CLL / SLL (352,353,2071,2265). They are included in the section on ‘Cutaneous involvement in primary extra-cutaneous B-cell lymphomas’ in the WHO classification of Haematolymphoid tumours. For each type of neoplasm a cell of origin is proposed.

**Histiocytic and dendritic cell neoplasms**
Histiocytic and dendritic cell neoplasms are derived from the phagocytic and accessory cells, which have a major role in the processing and presentation of antigen to lymphocytes and which are bone marrow derived. The origin of the B-antigen presenting follicular dendritic cells remains to be established. They are not of bone marrow origin; an origin from either a fixed stromal/mesenchymal cell or from blood vessel endothelium are the two favoured options (1068). Histiocytic and dendritic cell neoplasms are rare. Of these tumours, only Langerhans cell histiocytosis has a significant incidence of ear disease by virtue of involvement of the temporal bone and middle ear.
Langerhans cell histiocytosis

Definition
Langerhans cell histiocytosis (LCH) is a neoplastic proliferation of Langerhans cells, with expression of CD1a, S100 and the presence of Birbeck granules by ultrastructural examination.

ICD-O code 9751/1

Synonyms
Histiocytosis X, Langerhans cell granulomatosis. Clinical variants have been referred to as Letterer-Siwe disease, Hand-Schuller-Christian disease and solitary eosinophilic granuloma of bone.

Epidemiology
LCH is rare (1511). The incidence is about 5 per million with most cases occurring in children (1887). Bone involvement in LCH counts for less than 1% of all bone lesions. There is a wide distribution of age from a few months to the 9th decade of life (1066). Males are affected more often than females and the disease is more common in Whites of northern European origin than Blacks.

Etiology
This is unknown. There may be an association with a history of neonatal infection but there is no evidence of viral involvement (1672).

Localization
Three overlapping syndromes are recognised (1511). Unifocal disease occurs in the majority of patients and usually involves bone (solitary eosinophilic granuloma). It is the bones of skull which are particularly affected, followed in frequency by the femur, pelvic bones and ribs. Less commonly, unifocal disease is confined to a lymph node, skin or lung. LCH involving lungs in adults is nearly always associated with smoking and is thought to represent a different, possibly reactive disease (2684A). In multifocal, unisystem disease (Hand-Schuller-Christian disease), several sites in one organ, almost always bone, are affected.

In multifocal, multisystem disease (Letterer-Siwe disease) many organs are involved such as bones, skin, liver, spleen, lymph nodes and bone marrow. Any bone may be involved, with the highest frequency occurring in the bones of the skull in children (1511). In temporal bone disease the lesion involves the medial part of the external auditory meatus (2099).

Clinical features
Pain and swelling of the affected area is the most common presentation. In children with temporal bone involvement, the presenting features can simulate those of otitis media and mastoiditis because of otorrhea and mastoid and facial swelling. Radiologically, early lytic lesions can suggest an aggressive disease process. Mutifocal unisystem disease is usually confined to young children, and the multiple destructive bone lesions are often associated with adjacent soft tissue masses. With skull bone involvement there may also be exophthalmos and diabetes insipidus if the pituitary is affected and tooth loss, if the jaw bones are involved. Multifocal multisystem disease usually occurs in infants and in addition to bone lesions, there are fever, skin involvement, hepatosplenomegaly and pancytopenia because of bone marrow involvement.

Macroscopy
The involved tissue is soft and usually red. If haemorrhage and necrosis are present, the colour may be yellow due to the presence of lipid and many eosinophils.

Histopathology
Crucial to the diagnosis is the recognition of the Langerhans cell. It is the nuclear appearances which are so distinctive; the nuclei are folded or grooved resembling a coffee bean or lobulated and indented. The nuclear chromatin is finely dispersed, nucleoli are inconspicuous and the nuclear membranes are thin. Mitotic activity is quite variable; sometimes there are up to 6 mitoses per sq. mm. In bone lesions Langerhans cells are found in nests and clusters. Necrosis is often present and should not be interpreted as suggesting aggressive disease. Admixed with the Langerhans cells are eosinophils, sometimes in large numbers, lymphocytes, neutrophils and plasma cells. Multinucleated osteoclast-like giant cells and lipid laden foamy macrophages can usually be identified. In old lesions foamy macrophages are numerous and there is significant fibrosis. The appearances of the lesions are so characteristic that the diagnosis can be made on cytological preparations, including touch preparations.

Immunoprofile
Neoplastic Langerhans cells resemble normal Langerhans cells in their expression of CD1a (675) and S-100 protein, the latter being expressed in both nuclei and cytoplasm (1835). They also express CD4, vimentin, HLA DR and placental alkaline phosphatase (PLAP). CD68 and lysozyme are variably and only weakly expressed; and there is no expression of the follicular dendritic cell markers CD21 or CD35. Immunostaining for Ki67 shows a proliferation index of between 2% and 25%.

While their phenotype resembles that of normal Langerhans cells it is not identical, for in contrast their normal counterparts do not express placental alkaline phosphatase (PLAP) and also show differences in the expression of adhesion molecules (557).

Electron microscopy
As in normal Langerhans cells, neoplastic Langerhans cells contain the unique cytoplasmic organelle called the Birbeck or Langerhans granule. ‘Granule’ is something of a misnomer, since the characteristic shape is that of a tennis racket or long necked flask. These structures, which vary in length from 200-400 nanometres, are pentalaminar rods measuring 33 nanometres in width with a vesicular expansion at one end. They
arise from the cell membrane (1066). Their function is unknown.

Genetics
In all of the clinical syndromes/variants of LCH, studies of the X-linked androgen receptor gene have demonstrated that the proliferation of Langerhans cells is clonal (2796,2873).

Prognosis and predictive factors
It is the demonstration that Langerhans cell histiocytosis represents a clonal proliferation that has led to its acceptance as a neoplastic disorder. However, the prognosis for patients with either monostotic or limited polyostotic disease is good. Death is rare in LCH and is associated with disseminated forms of the disease. Thus the clinical outcome directly relates to the number of organs affected (946). The overall survival of patients with unifocal disease is 95% dropping to 75% with 2 organs involved and with further drops with increasing organ involvement. In about 10% of patients there is progression of unifocal lesions to multifocal disease. The presence of cytological atypia or an increased mitotic rate does not appear to correlate with prognosis (2181).
Secondary tumours

Definition
Neoplasms which originate from sites other than within the structures of the ear i.e. external auditory canal, middle ear and temporal bone. These may be metastatic via blood or lymphatic channels from non-contiguous sites, or spread directly from a contiguous site by invasion of surrounding tissues or extension along / through existing channels.

Epidemiology
Secondary neoplasms in the ear / temporal bone are rare, amounting to 5-6% of 2,528 benign and malignant ear neoplasms compiled from surgical pathology accessions from four institutions (549). Among the 1,781 ear neoplasms listed from the U.S. Armed Forces Institute of Pathology, only 31 (1.74%) had “metastasized” to the ear, most of these to the middle ear. However, post-mortem histologic studies of temporal bones reveal metastatic cancer in 47 (22%) of 212 cancer patients (895), virtually all of whom had had disseminated disease. The epidemiology of metastases involving the temporal bone is virtually identical to that cited for the skeletal system (775).

Incidence and mortality
Virtually all patients with metastases to the ear/temporal bone have a known primary malignancy elsewhere and widespread metastatic disease. Breast is by far the most common malignancy metastasizing to the temporal bone, followed by lung/bronchus, prostate, melanoma and thyroid. In spite of the frequency of metastases of renal carcinoma to bone, metastasis to the temporal bone appears to be rare, although a recent review reported renal origin in 9% of cases (1748).

However among 12 patients with occult carcinoma found incidentally at autopsy only one had distant metastases: prostate carcinoma with widespread bone metastases including temporal bone. Among 18 patients, “adequately treated” and clinically free of cancer, none had residual cancer at autopsy, including temporal bone (895).

Metastasis through direct extension
Direct extension into the ear / temporal bone occurs from the upper aero-digestive tract via the Eustachian tube and middle ear, and from the posterior fossa of the skull via the internal auditory canal. Direct invasion through bone and soft tissues occurs through the skull base and about the external ear in the parotid area. Head and neck primary tumours, excluding thyroid, account for the largest number (22%) of secondary tumours involving the ear by direct extension and/or invasion..

Localization
Blood borne metastases tend to localize to the petrous ridge (83%) and mastoid (28%) and are usually bilateral, multiple and associated with metastases to other bones. Contiguous spread via existing channels: anteriorly from the upper aerodigestive tract to the middle ear (21%) via the Eustachian tube (14.5%); posteriorly from the brain and meninges via the internal auditory meatus (28%) to the inner ear.

Invasion of bone and soft tissue with extension into the base of the skull and external auditory canal and middle ear and mastoid occurs with paragangliomas and malignancies of the parotid gland.

Clinical features
Metastases to the temporal bone generally occur late in the course of the disease. Signs and symptoms in 101 patients with a history of malignant neoplasm included hearing loss (28%), vertigo (10%) facial palsy (9%), tinnitus (7%), otalgia (5%), otorrhea (2%), external canal mass (2%), nystagmus (2), no symptoms (36%) (895,1863).

Histopathology
The secondary tumours maintain the phenotype of the primary. Since these secondary tumours occur in late stages of known malignant disease, evaluation of the unknown primary is rarely undertaken. Poorly differentiated malignancies invading the external auditory canal from a parotid lesion require consideration of parotid carcinoma, melanoma and adenocarcinoma of the canal. Biopsy of secondary temporal bone lesions, except for cerebellopontine angle tumours, is rarely undertaken and their origin is inferred by imaging studies, a history of malignancy or pathologic or cytologic studies of contiguous lesions.

Table 7.4 Site(s) of origin of tumours metastatic to temporal bone (895,1863). Age range, 2 - 87 years.

<table>
<thead>
<tr>
<th>Site</th>
<th>Total number (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Breast</td>
<td>27 (22%)</td>
</tr>
<tr>
<td>Head and neck incl. brain (7) and choroid plexus (1)</td>
<td>26 (21%)</td>
</tr>
<tr>
<td>Lung/bronchi</td>
<td>12 (10%)</td>
</tr>
<tr>
<td>Prostate</td>
<td>10 (8%)</td>
</tr>
<tr>
<td>Malignant melanoma</td>
<td>7 (6%)</td>
</tr>
<tr>
<td>Unknown origin</td>
<td>7 (6%)</td>
</tr>
<tr>
<td>Thyroid</td>
<td>5 (4%)</td>
</tr>
<tr>
<td>Miscellaneous other sites</td>
<td>9 (23%)</td>
</tr>
<tr>
<td>Total</td>
<td>123</td>
</tr>
</tbody>
</table>

Table 7.5 Temporal bone metastatic sites from 47 patients who died with malignancy University of Minnesota. From T.I. Gloria-Cruz et al. (895).

<table>
<thead>
<tr>
<th>Site</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Petrous apex</td>
<td>83%</td>
</tr>
<tr>
<td>Mastoid</td>
<td>27.6%</td>
</tr>
<tr>
<td>Internal auditory canal</td>
<td>27.6%</td>
</tr>
<tr>
<td>Middle ear</td>
<td>21.1%</td>
</tr>
<tr>
<td>Facial nerve</td>
<td>19.7%</td>
</tr>
<tr>
<td>Eustachian tube</td>
<td>14.5%</td>
</tr>
<tr>
<td>Otic capsule</td>
<td>13.2%</td>
</tr>
<tr>
<td>Cochlea</td>
<td>10.5%</td>
</tr>
<tr>
<td>Vestibule</td>
<td>9.2%</td>
</tr>
<tr>
<td>External ear</td>
<td>9.2%</td>
</tr>
<tr>
<td>Membranous labyrinth</td>
<td></td>
</tr>
<tr>
<td>Cochlear duct</td>
<td>7.9%</td>
</tr>
<tr>
<td>Saccule</td>
<td>3.9%</td>
</tr>
<tr>
<td>Utricle</td>
<td>3.9%</td>
</tr>
<tr>
<td>Semicircular canals</td>
<td>3.0%</td>
</tr>
</tbody>
</table>
CHAPTER 8

Tumours of the Paraganglionic System

The paraganglionic system develops early in gestation and is of neural crest origin. It consists of two components – the adrenal medulla and a diffuse collection of extra-adrenal paraganglia.

The extra-adrenal paraganglia are specialized collections of neuroendocrine cells that migrate in close association with the cranial nerves, large blood vessels, and autonomic nerves and ganglia. They vary in size from those that are just barely visible, such as the carotid bodies, to those that are apparent at the microscopic level, such as the laryngeal paraganglia.

As a group, neoplasms of the extra-adrenal paraganglia (paragangliomas) are uncommon and occur most often above the clavicles. Since paragangliomas are discussed in depth in the WHO Classification of Tumours of Endocrine Organs (577), they are only briefly summarized in this chapter.
Tumours of the paraganglionic system: Introduction

Introduction

The extra-adrenal paraganglia can be divided into sympathetic and parasympathetic types. Although they are indistinguishable at the cellular level, they differ in their anatomic distribution and secretory products. The sympathetic paraganglia are found primarily in the axial regions of the trunk along the prevertebral and paravertebral sympathetic chains and in connective tissue in or near pelvic organs. In contrast, parasympathetic paraganglia are localized almost exclusively in the head and neck along the branches of the glosopharyngeal and vagus nerves. Although both types of paraganglia produce catecholamines, clinical signs of excess production are usually associated with those that are of sympathetic origin. Tumours associated with significant amounts of epinephrine (adrenaline) are almost always of sympathetic origin. Those lesions that occur in patients with hypoxemia, in contrast, are typically parasympathetic in origin. Overlaps in their anatomic distribution and secretory expression, however, do occur. Parasympathetic paragangliomas, in contrast to their sympathetic counterparts, are also more often familial and less likely to be malignant. Paragangliomas of the head and neck are found primarily at the bifurcation of the common carotid artery, in the middle ear—temporal bone, along the course of the vagus nerve, and exceptionally, in the orbit, nasal cavity, paranasal sinuses, nasopharynx, larynx, trachea and thyroid.

Anatomy

Carotid body paraganglia

Carotid body paraganglia are paired, bilateral, usually symmetrical agglomerates of specialized neuroendocrine tissue located at the bifurcation of the common carotid artery along its posteromedial wall, either within or immediately external to the adventitia. They are anchored to the artery by a band of fibrovascular tissue referred to as the ligament of Mayer. Each measures about 3-7 mm in greatest dimension and weighs 3-15 mg [894, 1034, 2231, 2879]. They function as chemoreceptors sensitive to changes in oxygen tension, carbon dioxide content and hydrogen ion concentration.

Jugulotympanic paraganglia

Small paraganglia with a structure similar to the carotid body have been described in the ear [957]. More than 50% of these structures are situated in relation to the jugular bulb; a minority are found under the mucosa of the middle ear in the region of the medial promontory wall. The tumours arising from these paraganglia form the more frequent jugular paraganglioma (glomus jugulare) and the less frequent tympanic paraganglioma (glomus tympanicum), respectively.

Vagal paraganglia

The vagus nerve (from the Latin “vagus” meaning wandering and meandering) arises from 8-10 rootlets on the lateral border of the medulla that converge to form a cord on entering the jugular foramen. It is the longest cranial nerve and has a superior ganglion, which lies within the jugular foramen, and just below this, a middle ganglion. A third, much larger ganglion, known as the ganglion nodosum, lies more inferiorly. It is approximately 2.5 cm long and lies high in the neck, just behind the internal carotid artery. Vagal paraganglia do not form a discrete “body” as seen at the carotid artery bifurcation, but rather consist of 6-7 small, dispersed agglomerates of paragangliomous tissue. They may be found within (intravagal), or adjacent to (juxtavagal), the vagus nerve, usually at the level of the nodose ganglion. In rare instances, paragangliomous tissue may be found in sites just above or below the nodose ganglion. Their function is unknown, but they may have a role as a chemoreceptor and moderator of the cardiopulmonary system.

Laryngeal paraganglia

The larynx contains two pairs of paraganglia that are divided into two groups: superior and inferior. The superior paraganglia are between 0.1 mm and 0.3 mm in diameter. They occur bilaterally in the upper one-third of the false cord, adjacent to the superior margin of the thyroid cartilage, in close proximity to the superior laryngeal artery and nerve [2731]. The inferior paraganglia are also bilateral but larger than the superior group, averaging 0.3-0.4 mm in diameter [1331].
**Table 8.4** Differential diagnosis of paraganglioma of the head and neck.

<table>
<thead>
<tr>
<th>Stain</th>
<th>Paraganglioma</th>
<th>Carcinoid</th>
<th>Medullary Thyroid Carcinoma</th>
<th>Anaplastic Carcinoma</th>
<th>Melanoma</th>
<th>Renal Cell Carcinoma</th>
</tr>
</thead>
<tbody>
<tr>
<td>Synaptophysin</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Keratin</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>HMB-45</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Renal cell carcinoma antigen</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>+</td>
</tr>
<tr>
<td>Calcitonin</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Thyroid transcription factor</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Congo red (amyloid)</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>+</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

They characteristically occur between the inferior border of the thyroid cartilage and cricoid cartilage or between the cricoid cartilage and first tracheal ring in relationship to the inferior laryngeal artery and nerve. Aberrant and accessory paraganglia are described [1331,2878]. Occasionally, paraganglia may even be found within nerves rather than adjacent to them. This is especially so regarding the recurrent laryngeal nerve [525]. The physiological role of the laryngeal paraganglia is also unknown. They may possibly serve as extracarotid chemoreceptors or have some effect on respiration via the larynx.

**Terminology**
The preferred terminology for tumours of the extradrenal paraganglia is “paraganglioma”, prefixed by the anatomic site of origin, for instance, carotid body paraganglioma [1412]. If the tumour is functional or malignant, it would be designated, for example, as a functional carotid body paraganglioma or a malignant carotid body paraganglioma. Although the adrenal medulla is technically a paraganglion, a tumour arising from this site is still regarded as a phaeochromocytoma rather than a paraganglioma.

**Diagnostic procedures**
Computed tomography (CT) with contrast medium and magnetic resonance imaging (MRI) with gadolinium are invaluable in defining the location, size and extent of a paraganglioma [1587,2124]. The typical CT appearance of a carotid body and vagal paraganglioma is that of a homogenous, hypervascular, well-defined soft tissue mass. If there has been haemorrhage or focal thrombosis, a heterogenous pattern of enhancement will be seen. CT scans of jugulotympanic paragangliomas may also show expansion and erosion of the jugular foramen. As the tumour expands, it may destroy the surrounding bony labyrinth and ossicular chain and extend into the region of the cerebellopontine angle [2124].

MRI characteristics of all paragangliomas are similar. A well-defined hypointense mass with areas of signal void is typically seen on T1-weighted images [1587]. A “salt and pepper” pattern is commonly seen in all lesions larger than 2 cm. This pattern is usually seen on T2-weighted images and is due to areas of high vascularity associated with haemorrhage or slow blood perfusion. In addition to providing superior definition, MRI can also detect much smaller paragangliomas than CT scans. Octreotide scintigraphy is an important adjunct. It can be used not only to confirm the diagnosis of a neuroendocrine neoplasm, but also to detect additional occult tumours, to separate postoperative changes from residual or recurrent disease and for screening patients at risk for familial paragangliomas [1398,1587]. Ultrasound has a limited role in the evaluation of head and neck paragangliomas. It is useful in the evaluation and follow-up of carotid body paragangliomas and to some extent vagal paragangliomas. Its use in the detection and assessment of jugulotympanic paragangliomas is limited because of the surrounding bone. Although non-invasive imaging has almost universally replaced angiography as the primary radiographic procedure for diagnosing paragangliomas, angiography still remains an important component in the management of these patients, especially in regards to preoperative embolization to reduce the blood supply of the tumour.

**Genetics**
It is commonly stated that about 10% of all paragangliomas of the head and neck are familial and inherited as an autosomal dominant trait with genomic imprinting [951,1670,2665]. There is no tumour when the gene is inherited from the mother. Paternal transmission of the gene, however, results in tumour(s) in children even if the father is clinically unaffected. Some investigators are of the opinion that because of skipping of generations after maternal transmission of the gene, that the incidence of familial paragangliomas is vastly underestimated and may be as high as 50% of all cases [2678]. Genetic linkage analyses of several large families with hereditary paragangliomas have identified three loci associated with these tumours – paraganglioma 1 (PGL1) on chromosome 11q23, PGL2 on chromosome 11q13.1 and PGL3 on chromosome 1q21-q23 [1080,1634,1897]. Studies have identified the PGL1 gene as SDHD and the PGL3 gene as SDHC, both of which encode mitochondrial respiratory chain proteins [181,182,889]. In a study of 8 patients with sporadic (non-familial) paragangliomas of the head and neck, 3 exhibited deletions at chromosome 11q, 2 at 11q22-23 and 1 at 11q13 [213]. This suggests that sporadic and familial paragangliomas may share a similar molecular-genetic pathogenesis. It is now possible through genetic analysis to identify early on those patients who are at risk for familial paragangliomas, with the possibility of gene therapy on the horizon [2025].
Introduction
A neuroendocrine neoplasm derived from carotid body paraganglia composed of chief and sustentacular cells arranged in a characteristic (Zellballen) pattern.

ICD-O code 8692/1

Synonyms
Carotid body tumour, chemodectoma, glomus tumour, non-chromaffin paraganglioma, neuroendocrine tumour.

Age and sex distribution
They occur primarily in adults averaging 40-50 years of age and are rare in children. At sea level, the sex distribution ranges from 1:1-1:4 in favour of females {136,951,1410,1869,2155}. However, at altitudes above 2000 meters, there is a female predominance of 8.3:1 {2188}. It has been postulated that the monthly loss of blood through menstruation in women and a larger pulmonary capacity and greater enthusiasm for sports and athletic conditioning in men may allow males to escape chronic hypoxia and account for this wide gender gap {2188}.

Etiology
Familial inheritance and chronic hypoxia are the only known risk factors.

Localization
The tumours occur at the bifurcation of the common carotid artery with no significant lateralization to either side of the neck. As they enlarge, they may become adherent, invade or incorporate the external and/or internal carotid arteries.

Clinical features
Although uncommon, carotid body paragangliomas are regarded as the most common paraganglioma of the head and neck.

Signs and symptoms
The typical presentation is that of a slowly enlarging, asymptomatic mass deep to the anterior border of the sternocleidomastoid muscle just below the angle of the mandible. The tumour can be moved from side to side but with little or no movement in a vertical plane (Fontaine's sign). Occasionally, it may be associated with pain, hoarseness, dysphagia, Horner's syndrome, headache, syncope, bruit or thrill. Functional tumours with catecholamine - induced hypertension are exceptional {509}.

Infrequently, the tumour may be bilateral or associated with paragangliomas in other sites (usually a vagal or jugulotympanic paraganglioma) or occur in combination with a phaeochromocytoma, a well-differentiated thyroid carcinoma or a component of multiple endocrine neoplasia syndrome or Carney triad {481,509,1391,1422,1430}. Hereditary deficiencies of clotting factors VII and X have also been observed in a few patients with familial carotid body paragangliomas {1376}.

Histopathology
These highly vascular tumours are composed of two cell types, chief and sustentacular, arranged in a characteristic alveolar or Zellballen pattern. The chief cells (type I cells, epithelioid cells) are more numerous and contain catecholamine-bound neurosecretory granules as seen ultrastructurally. The sustentacular cells (type II cells, supporting cells) are devoid of neurosecretory granules and are characteristically located at the periphery of the Zellballen. The chief cells often show considerable nuclear enlargement and hyperchromatism and contain cytoplasm that varies from pink to clear, to amphophilic and which may be vacuolated. Spindle-shaped chief cells are uncommon and mitoses are sparse to absent. Vascular and perineural invasion are infrequent and have no prognostic significance.

Immunoprofile
The chief cells express synaptophysin,
chromogranin and neuron-specific enolase. They are negative for cytokeratin, carcinoembryonic antigen, S-100 protein and calcitonin. The sustentacular cells express S-100 protein and glial fibrillary acidic protein.

Malignant carotid body paraganglioma
Paragangliomas are divided into non-invasive (circumscribed, encapsulated), locally invasive and metastatic categories. Some locally invasive tumours may even cause the death of the patient. Although clinically malignant, these tumours are still regarded as locally invasive. A tumour is considered malignant only if there is metastasis to regional lymph nodes or to more distant sites, such as the lungs and bones. The incidence of malignant (metastasizing) carotid body paragangliomas ranges from 2-13% [136]. Unfortunately, the clinical behaviour cannot be predicted on the basis of routine histology. Such features as nuclear pleomorphism, mitotic activity, necrosis and vascular – perineural invasion are unreliable prognosticators and have been found in benign as well as malignant tumours [136]. Other findings such as DNA ploidy, absence of sustentacular cells, number of expressed neuropeptides, assessment of argyrophilic nucleolar organizer regions and proliferative markers (PNCA, Ki-67) show no consistent correlation with histological behaviour [661,871,1335,1336, 1382, 1526,2678,2748,136].

Sporadic (non-familial) carotid body paragangliomas are more likely to be malignant than those that are familial – 12% versus 2.5% [951]. Metastases may be apparent at the time of initial therapy or may not become apparent until 20 years later [1967]. Surgery with or without adjuvant irradiation is used for local disease. Chemotherapy, however, is largely ineffective [1656]. The overall 5-year relative survival is 59.5%. If the metastases are confined to regional lymph nodes, the 5-year survival is 76.8% but decreases to 11.8% for patients with distant metastases [1451].

Differential diagnosis
The differential diagnosis includes carcinoid, medullary thyroid carcinoma, anaplastic carcinoma, metastatic melanoma and renal cell carcinoma. Immunostains are useful in separating these tumours.

Genetics
One-third of familial carotid body paragangliomas are bilateral, as opposed to 4% bilateral sporadic (non-familial) cases [951]. These may appear synchronously or asynchronously. C-myc, bcl-2 and c-jun are abnormally expressed in some tumours and may contribute to tumorigenesis [2719, 2721]. Overexpression of TP53, however, does not appear to be an etiologic factor [2720].

Prognosis and predictive factors
Carotid body paragangliomas are slowly growing tumours with a median growth of 0.83 mm per year and a median doubling time of 7.13 years [1205]. Surgery, often with sacrifice of one or more branches of the carotid arterial system, is the treatment of choice. Somewhere between 0-10% of tumours will recur. This is not necessarily a sign of malignancy but rather inadequate excision and regrowth.
Jugulotympanic paraganglioma

Definition
A neoplasm arising from one or other of the paraganglia situated in the vicinity of the jugular bulb or on the medial promontory wall of the middle ear.

ICD-O code 8690/1

Synonyms
Jugulotympanic chemodectoma, glomus jugulare tumour, jugular glomus tumour, glomus tympanicum tumour, tympanic glomus tumour.

Epidemiology
Solitary jugulotympanic paragangliomas arise predominantly in females. The age range is between 13 and 85 years with a mean age of about 50 years. The familial type occurs predominantly in men.

Localization
Most jugulotympanic paragangliomas arise from the paraganglion situated in the wall of the jugular bulb. A minority arise from the paraganglion situated near the middle ear surface of the promontory. The distinction between jugular and tympanic paragangliomas can easily be made in the patient by modern imaging methods with which the jugular neoplasm is identified as arising from the jugular bulb region and shows evidence of invasion of the petrous bone, while the tympanic neoplasm is confined to the middle ear. Jugulotympanic paragangliomas may also be multicentric or coexist with tumours of other types. They may be bilateral in the same patient and coexist with carotid body paragangliomas which may also be bilateral [1949]. They may also coexist with adrenal gland pheochromocytomas.

Clinical features
Most patients present with conductive hearing loss. Pain in the ear, facial palsy, haemorrhage, and tinnitus are also described. On examination, a red vascular mass is seen either behind the intact tympanic membrane or sprouting through the latter into the external canal. Surgical approach to the mass at biopsy often results in severe bleeding.

Etiology
The etiology of the solitary type is unknown. In the multiple familial type there is evidence of a gene mutation on chromosome 11.

Macroscopy
The neoplasm is an irregular reddish mass. In the jugular variety, the petrous temporal bone and the middle ear space are largely replaced by red, firm material as far as the tympanic membrane. The otic capsule is rarely invaded by paraganglioma. Investigation of a paraganglioma in an autopsy temporal bone by the microslicing method, showed the shape of the jugular bulb to be retained, but the lumen was completely filled by neoplasm. The tumour invaded the apical region of the petrous temporal bone and the middle ear space was completely filled by neoplasm as far as the tympanic membrane. However, the surgical specimen is usually fragmented.

Histopathology
The histological appearances of the jugular and tympanic paragangliomas are similar, resembling that of the carotid body paraganglioma. Epithelioid, small, uniform cells, with finely granular cytoplasm are separated by numerous blood vessels. The tumour cells often form clusters or “Zellballen” with peripheral flattened cells. Nuclei are usually uniform and small, but diagnosis is sometimes made difficult by the presence of bizarre or multinucleate cells which, however, do not indicate malignancy. A prominent fibrous stroma is sometimes present.

Fig. 8.4 Jugulotympanic paragangliomas. A Axial CT scan of petrous bone. On the right side, there is erosion of the cortex of the jugular foramen in keeping with a jugular paraganglioma (upper arrow). Note the normal jugular foramen (lower arrow) on the left. B Axial CT scan of petrous bone. Soft tissue mass in the posterior hypotympanum (upper arrow). The adjacent permeative erosion (lower arrow) suggests that this is a tympanic paraganglioma.
Immunoprofile
The immunoprofile of these tumours has been covered in an earlier section (Immunoprofile p. 364-365).

Electron microscopy
Paragangliomas shows membrane-bound, electron-dense neurosecretory granules in the cytoplasm of the tumour cells consistent with catecholamine content [2277].

Prognosis and predictive factors
Jugulotympanic paraganglioma is a neoplasm of slow growth. The jugular variety infiltrates the petrous bone, but distant metastasis is rare. Radiation therapy, and in some cases surgery, offers a high rate of cure for the localized neoplasms.

Fig. 8.5 Jugulotympanic paragangliomas. A Section of autopsy case of jugulotympanic paraganglioma showing tumour behind tympanic membrane. B Histological appearance of jugulotympanic paraganglioma. C Jugulotympanic paraganglioma showing numerous sustentacular cells (S100 protein). D Jugulotympanic paraganglioma showing bizarre cells some of which are multinucleate.
Vagal paraganglioma

Definition
A neuroendocrine neoplasm derived from paraganglia found within or adjacent to the vagus nerve usually in the vicinity of the ganglion nodosum.

ICD-O code 8693/1

Synonyms
Vagal body paraganglioma, glomus tumour, glomus vagale, chemodectoma, non-chromaffin paraganglioma, neuroendocrine tumour.

Epidemiology
Age and Sex distribution
Vagal body paragangliomas are more common in women (64%) and occur over a broad age range (19-86 years) with an average of 45-55 years [215,282,689,1410,1411,1736,1868,2659].

Etiology
Most are sporadic. Some are related to familial inheritance. Although chronic hypoxia may lead to hyperplasia of vagal paraganglia, there is no conclusive evidence, in contrast to the carotid body paraganglioma, that it leads to the development of a vagal paraganglioma [1409].

Localization
The tumours typically occur in the rostral portion of the vagus nerve in the vicinity of the ganglion nodosum. In a review of 99 vagal paragangliomas in which the side of origin was indicated, 56% arose on the right side of the neck, 39% on the left side and 5% were bilateral [2879].

Clinical features
The vagal paraganglioma is the third most frequent paraganglioma of the head and neck, exceeded in frequency only by the carotid body and jugulotympanic paragangliomas. It characteristically presents as a slowly enlarging, asymptomatic mass at the angle of the mandible and/or as a bulge in the lateral oropharyngeal wall. As it increases in size, deficits of cranial nerves IX, X, XI and XII and the cervical sympathetic chain are common, resulting in unilateral vocal cord dysfunction, dysphagia, atrophy of the tongue, shoulder weakness and Horner syndrome. At the time of diagnosis, anywhere from 35-65% of patients may manifest one or more cranial nerve deficits.

Functional tumours with catecholamine-induced hypertension are distinctly uncommon, occurring in only 3.6% of all tumours.

Imaging
Vagal paragangliomas are highly vascular and are located in the suprahyoid neck, well above the level of the carotid bifurcation and typically displace both external and internal carotid arteries anteromedially.

Macroscopy
The tumours are fusiform or circular and abut directly onto the base of the skull. They usually range from 2.0-6.0 cm and are firm, rubbery, well circumscribed and surrounded by a thin, sometimes locally thickened fibrous capsule. In a few instances, they may be poorly defined and locally infiltrative. On cut surface, they have a variegated yellow, tan, pink, red or brown appearance with fibrosis and haemorrhage or they may be uniformly homogenous. A portion of one or more large nerves, usually the vagus, is often attached.

Histopathology
The histopathology, immunoprofile, ultrastructural features and differential diagnosis are similar to those previously

Table 8.5 Vagal paraganglioma: clinico-pathologic features*

<table>
<thead>
<tr>
<th>Feature</th>
<th>Data</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average age (n=139)</td>
<td>45 – 55 years (range 19 – 86 years)</td>
</tr>
<tr>
<td>Gender (n=139)</td>
<td>64% female (range 50 – 85%)</td>
</tr>
<tr>
<td>Multicentric tumours (n=124)</td>
<td>33% (range 10 – 46%)</td>
</tr>
<tr>
<td>Familial history (n=124)</td>
<td>17% (range 0 – 47%)</td>
</tr>
<tr>
<td>Functional tumours (n=139)</td>
<td>3.6% (range 0 – 11%)</td>
</tr>
<tr>
<td>Local recurrence (n=126)</td>
<td>8% (range 0 – 20%)</td>
</tr>
<tr>
<td>Malignant tumours (n=139)</td>
<td>7% (range 0 – 16%)</td>
</tr>
</tbody>
</table>

*Data based on references [215,282,689,1410,1411,1736,1868,2659].

Fig. 8.6 Vagal paraganglioma. A Note the expanded portion of the vagus nerve which represents the neoplasm. A small group of lymph nodes is attached at the center of the specimen. B Cross section shows a spongy, focal haemorrhagic tan surface. A segment of vagus nerve is attached.
described for the carotid body paraganglioma.

**Malignant vagal paraganglioma**

Overall 7% of vagal paragangliomas are malignant by virtue of metastases. In one review of 15 malignant vagal paragangliomas, 73% were associated with cervical lymph node metastasis and 27% with distant metastases (lung, bone, liver and brain) \{1050\}. Most metastases are apparent either at or within four years of diagnosis.

**Genetics**

Patients with sporadic (non-familial) vagal paragangliomas may have more than one paraganglioma and should always be evaluated for this possibility. The incidence of finding multiple tumours in this population varies according to the thoroughness of the examination and the length of follow-up. When multiple, the tumours may appear synchronously or asynchronously. In a collective review of 124 vagal paragangliomas, 33% were associated, either at the time of diagnosis or on follow-up, with additional paragangliomas, usually a carotid body or, less frequently, a jugulotympanic paraganglioma. Seventeen percent of patients also had a family history of paragangliomas.

DNA analysis of 10 vagal paragangliomas utilizing image analysis, revealed 5 to be diploid, 4 diploid – tetraploid and 1 aneuploid \{137\}. DNA abnormalities are, therefore, common in these tumours and cannot be used to predict prognosis.

**Prognosis and predictive factors**

Vagal paragangliomas are slowly growing with an estimated median growth rate of one millimetre per year and a median doubling time of 8.89 years \{1205\}. Options for treatment include surgical resection, radiation, and, in selected cases due to their slow growth rate, even observation. Most clinicians favour surgery. Almost invariably, the vagus nerve and sometimes even other cranial nerves, have to be sacrificed. In those instances where the nerve is preserved, function typically remains permanently impaired. Failure to remove the nerve may also predispose the patient for future recurrence.

Radiation is used for elderly patients who are poor operative risks or for those unfortunate individuals who have bilateral vagal paragangliomas (the larger tumour is preferentially excised while the smaller one is irradiated). Following surgery, 8% of vagal paragangliomas will develop local recurrence. The tumour may recur as early as 12 months after therapy or as late as 22 years. Local recurrence is not necessarily a sign of malignancy but often results from inadequate excision.
Laryngeal paraganglioma

Definition
A neuroendocrine neoplasm derived from either the superior or inferior paraganglia of the larynx composed of chief and sustentacular cells arranged in a characteristic organoid (Zellballen) pattern.

ICD-O code
8693/1

Synonyms
Glomus tumour, chemodectoma, non-chromaffin paraganglioma, neuroendocrine tumour.

Age and sex distribution
Laryngeal paragangliomas are rare, with only 62 cases identified in a critical review of the world literature in 1994 [125,739]. They are three times more common in women and have been described in patients from 5-83 years of age (median 44 years) [739,2586].

Etiology
Other than a familial inheritance, there are no known risk factors.

Localization
The vast majority (82%) occur in the supraglottic larynx, presumably arising from the superior pair of laryngeal paraganglia, and present as a submucosal mass in the region of the aryepiglottic fold – false vocal cord [125]. Only 15% occur in the subglottis and 3% in the glottis. The right side of the larynx is more often involved than the left by a ratio of 2.3:1 [125].

Clinical features
Most patients present with more than one complaint and have been symptomatic for an average of 26 months (median 23 months; range 3 weeks to 12 years) [125]. The major symptom, by far, is hoarseness. Others, in decreasing order of frequency, are dysphagia, dyspnea, stridor, dysphonia, sore-painful throat, neck mass, haemoptysis, coughing, shortness of breath, foreign body sensation in the throat, and otalgia. Bruits and pulsation are usually absent.

Fig. 8.7 Diagram of the larynx showing location of laryngeal paraganglia.

Functional laryngeal paragangliomas are exceptional with only one possible case report [1438]. The patient, a 25-year-old woman with a supraglottic paraganglioma had sinus tachycardia and hypertension, which disappeared following surgical removal of the tumour. Neither the patient nor the tumor was evaluated for hormone production. Other alleged functional tumours that have been reported are probably atypical carcinoids [1184,1240,2691].

Imaging
Very few laryngeal paragangliomas have been evaluated preoperatively with angiograms. Of those that have, the blood supply of supraglottic paragangliomas has been from the superior thyroid artery, superior laryngeal artery or a branch of the external carotid artery [125,149,1123,1357,2236]. An angiogram of a single case of a subglottic paraganglioma showed the blood supply was via the thyrocervical trunk [1940].

Macroscopy
The tumours characteristically present as a well-circumscribed, tan, brown or reddish-brown 0.5-6.0 cm (average 2.6 cm) submucosal mass [125]. Cut surface varies from smooth to multinodular, with or without areas of fibrosis.

Histopathology
Histopathology, immunoprofile, ultrastructural features and differential diagnosis are similar to those of the carotid body paraganglioma.

Malignant laryngeal paraganglioma
Although commonly stated that 25% of all laryngeal paragangliomas are malignant [851,2413,2779], a critical review of these cases has revealed that almost all of these are examples of atypical carcinoids incorrectly labelled as malignant paragangliomas [125]. Current studies indicate that only about 2% of all laryngeal paragangliomas are malignant [739]. Only a single acceptable case has been reported. This involved a 36 year-old woman with a paraganglioma of the larynx that metastasized to the lumbar spine 16 years after the diagnosis [96,2212].

Genetics
Laryngeal paragangliomas may be associated with paragangliomas in other sites [125]. The most frequent association is with a carotid body paraganglioma [1020,2236,2679]. Cases have also been described associated with a jugulotympanic and a tracheal paraganglioma [510,1328]. Another example has been reported in a 35 year-old man with a family history of carotid body paragangliomas who presented with a subglottic paraganglioma [254]. DNA analysis of two laryngeal paragangliomas has shown both to be diploid [125].

Prognosis and predictive factors
Surgery is the treatment of choice, preferably through an external approach [739]. Endoscopic excision should be avoided (even for small lesions) because bleeding, which may be diffuse, may be difficult to control. Preoperative angiography and embolization, in an attempt to devascularize the tumour, is not essential since the superior thyroid artery can easily be ligated prior to resection. An elective neck dissection is not warranted. Seventeen per cent of patients have developed local recurrence from 1-16 years after initial excision [125].
Contributors
Source of charts and photographs

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Pathology and Genetics of Head and Neck Tumours is the latest volume in the new WHO series on histological and genetic typing of human tumours. This authoritative, concise reference book provides an international standard for pathologists and oncologists and will serve as an indispensable guide for use in the design of studies monitoring response to therapy and clinical outcome.

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