Chapter from the book *Carcinogenesis, Diagnosis, and Molecular Targeted Treatment for Nasopharyngeal Carcinoma*

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Imaging of Nasopharyngeal Carcinoma

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1. Introduction

1.1 Imaging of nasopharyngeal carcinoma

Nasopharyngeal carcinoma (NPC) is the most common neoplasm to affect the nasopharynx (NP). Arising from the epithelial lining of the nasopharyngeal mucosa, NPC is distinct from squamous cell carcinoma affecting other sites of the pharyngeal space. NPC typically originates in the lateral wall of the nasopharynx and is noted as a locally aggressive neoplasm with a high incidence of metastases to cervical lymph nodes. The primary tumour can extend within the nasopharynx and/or to the base of the skull, palate, nasal cavity or oropharynx. Distant metastases can arise in bone, lung, mediastinum, and, more rarely, the liver (Brennan, 2006).

The etiology of NPC is multifactorial and involves many environmental and genetic risk factors (Henderson et al., 1976). In particular, diets high in salt-preserved foods - such as salted fish, meat, eggs, fruits and vegetables in the Southern Asian diet - have been identified as possible causative factors acting through the carcinogen, N-nitrosodimethylamine (Yu et al., 1988). In contrast, frequent consumption of fresh fruits and/or vegetables - especially during childhood - is associated with a lower risk of NPC. In addition, studies have indicated a causal role for the Epstein-Barr virus (EBV) in the development of NPC (Chang & Adami, 2006). EBV is a DNA virus responsible for infectious mononucleosis, post-transplantation lymphoproliferative disease, and Burkitt’s lymphoma. Elevated IgG and IgA antibody titres against viral capsid antigen, early antigen, and latent viral nuclear antigens have been noted in NPC patients. Furthermore, these antibodies have been shown to precede tumour development by several years. EBV DNA, RNA, and gene products have also been detected in tumour cells (Chang & Adami, 2006). EBV serology is currently used as a screening tool for high-risk populations in southern China (Glastonbury, 2007). Certain genetic haplotypes – HLA A2, Bsn 2, B46, and B58 – are also associated with
increased risk (Ren & Chan, 1996). Other exposures implicated in NPC include cigarette smoking, other smoke, and occupational exposures including wood dust and industrial heat (Yu et al., 2010).

2. Epidemiology of NPC

NPC is a rare malignancy that exhibits a distinct ethnic and geographic variation (Chang & Adami, 2006). While in most regions of the world, age standardized incidence rates for both males and females are <1 per 100,000 person-years (Chang & Adami, 2006; Chong, 2006), higher rates are observed in a few well-defined populations. Studies have demonstrated intermediate incidence rates in several indigenous populations in Southeast Asia, the Arctic, and the Middle East/North Africa (ranging from 0.5 to 31.5 per 100,000 person-years in males and 0.1 to 11.8 person-years in females) (Chang & Adami, 2006; Parkin et al., 2002), and endemic rates in the natives of southern China reaching 20-30 per 100,000 person-years and 15-20 per 100,000 person-years amongst males and females, respectively, in the province of Guangdong (Ho, 1978; Yoshizaki et al., 2011). In addition, within geographic regions, a distinct ethnic variation also exists. For example, in the US, rates are highest amongst Chinese Americans, followed by Filipino Americans, Japanese Americans, Blacks, Hispanics, and finally Caucasians (Burt et al., 1992).

Globally, there are more than 80,000 incident cases and 50,000 deaths annually due to NPC (Parkin et al., 2005). NPC shows a male predilection of 2-3:1 (Parkin et al., 2002). In low risk populations, incidence increases with age, while in high risk populations, incidence peaks in the 6th decade of life and declines thereafter. NPC is rare in pediatric populations. Thus, the highest incidence rates are observed in middle-aged men (Chang & Adami, 2006).

3. Histopathology of NPC

Three histologic subtypes are recognized by the World Health Organization (WHO) classification system:

- Type 1 – squamous cell carcinoma; typically found in older adult populations
  - Type 1 is further subdivided into poorly, moderately, and well-differentiated
- Type 2 – non-keratinizing carcinoma
- Type 3 – undifferentiated or poorly differentiated carcinoma, including lymphoepithelioma and anaplastic variants; comprises most cases of childhood and adolescent NPC

In contrast to Type 1, Type 2 and 3 are associated with elevated EBV titres. In addition, Type 2 and 3 may also be accompanied by lymphoepithelioma, which is an inflammatory infiltrate of lymphocytes, plasma cells, and eosinophils. Histopathologic typing has prognostic significance, as Type 2 and Type 3 exhibit a more favourable prognosis and greater radiosensitivity than Type 1. Risk factors for Type 1 include cigarette smoke and ethanol consumption, whereas Type 2 and 3 are the endemic forms (Brennan, 2006).

In NPC, two histological patterns may be recognized: (1) the Regaud type which is a well-defined collection of epithelial cells surrounded by lymphocytes and connective tissue, and (2) the Schmincke type, which appears as diffuse tumours intermingled with inflammatory cells (Brennan, 2006).
4. Presentation of NPC

While some patients are asymptomatic, 50% to 70% of patients initially present with cervical lymphadenopathy in the form of a neck mass and are diagnosed by lymph node biopsy (Glastonbury, 2007). Symptoms at presentation may include trismus, pain, otitis media due to Eustachian tube dysfunction, nasal regurgitation due to paresis of the soft palate, and hearing loss. Depending on the degree of local infiltration, patients may also suffer from cranial nerve dysfunction and headache. Cranial nerve neuropathies (most often CN5 and CN6) are an indication of skull base infiltration. Larger growths may also produce nasal obstruction or bleeding and a "nasal twang". Metastatic spread may result in bone pain or organ dysfunction. Rarely, a paraneoplastic syndrome of osteoarthropathy may occur with widespread disease.

5. Role of imaging in the management of NPC

All stages of NPC patient management, from diagnosis and staging to treatment and follow-up, involve imaging. On presentation, a full diagnostic work-up for NPC involves a physical examination (including neurological examination of the cranial nerves), laboratory testing (including a complete blood count, liver function tests, and alkaline phosphate levels), an endoscopic-guided biopsy, as well as imaging studies. Since 6% of NPC is submucosal and cannot be seen on endoscopy (King et al., 2006), cross-sectional imaging studies, such as magnetic resonance imaging (MRI) or computed tomography (CT), are required to help confirm the diagnosis, as well as accurately demarcate the exact limits of pharyngeal wall involvement and tumour invasion into surrounding structures. Currently, MRI and CT are not routinely used for screening purposes; however, the radiologist should consider NPC whenever head and neck imaging is obtained, especially in high-risk patients – such as those of Asian descent – being evaluated for otitis media or with incidental findings of middle ear opacification.

Early findings of NPC on imaging include asymmetry of the nasopharynx and an obstructed Eustachian tube (ET) (Figure 1) (Glastonbury, 2007). Most NPC masses originate in the fossa of Rosenmuller, otherwise known as the lateral pharyngeal recess. Furthermore, involvement of the lateral pharyngeal recess may cause dysfunction of the ET – either directly or indirectly by infiltrating the surrounding musculature – leading to stasis of middle ear secretions and unilateral hearing loss (Figure 2). In cases of head and neck imaging for neck masses of “unknown primary,” careful attention should be paid to the possibility of NPC since cervical lymphadenopathy is the most common presentation.

Like other neoplasms, a mainstay in NPC treatment is staging. Staging contributes information regarding prognosis and helps guide treatment planning, facilitate stratification of treatment, and coordinate clinical studies (Mao et al., 2009; Yu et al., 2010). While in the past, CT was preferred, MRI is currently the imaging modality of choice for NPC staging using the American Joint Committee on Cancer (AJCC) tumour, node, metastasis (TNM) staging system (King et al., 1999; Liang, 2009; Ng et al., 1997). CT still has a role in the assessment of bony skull base involvement (Olmi, 1995), but MRI is considered superior to CT for assessing primary tumour invasion into surrounding soft tissue and bony structures, pharyngobasilar fascia invasion, invasion into the sinus of Morgagni, skull base invasion, as well as cavernous sinus extension and perineural disease (Liao et al., 2008; Sakata, 1999).
Fig. 1. Axial T2 weighted image shows a mass filling the left fossa of Rosenmuller (*).

Fig. 2. Axial T2 weighted image shows a right sided NPC that is resulting in right Eustachian tube dysfunction and fluid build up in the right mastoid.
MRI is also more reliable for differentiating between the primary tumor and retropharyngeal adenopathy (Chang, 2005; Chong, 1996; Chung, 2004; King, 2000). For patients with clinical or biochemical evidence of advanced disease, additional investigations – including bone scintigraphy, chest x-ray, CT of thorax, abdomen, and pelvis, and/or fluorodeoxyglucose (FDG)-positron emission tomography (PET)-CT imaging – may be indicated (Caglar, 2003; Chen et al., 2006; Chiesa & De Paoli, 2001; Chua et al., 2009; Comoretto et al., 2008; Lee, 1992).

The mainstay of treatment in NPC is radiation therapy (RT) with the addition of concurrent and/or adjuvant chemotherapy reserved for higher stages of disease. Surgical salvage (nasopharyngectomy) has also been used in cases of recurrent disease (Yu et al., 2010). The goal of treatment is to be curative. To be effective, RT must target the primary lesion as well as any associated neck adenopathy. Thus, cross-sectional imaging, in particular MRI, is needed to determine areas of disease infiltration. NPC has been shown to spread in a step-wise pattern along privileged pathways, such as the neuroforamina (Liang et al., 2009). In addition, involvement of critical structures located near the NP – including the cavernous sinus, pituitary gland, orbit and brainstem – must be evaluated to guide appropriate treatment and to spare these structures of the radiation dose if they are clear of disease (Yu et al., 2010). In the case of intensity-modulated RT (IMRT), CT is also required to correctly calculate dose based on absorption rates (Glastonbury, 2007).

Follow-up evaluation involves a baseline imaging study that is typically performed 2 to 3 months after completion of radiation treatment, followed by imaging every 3 to 6 months for the first 2 post-treatment years (Glastonbury, 2007). Any soft tissue signal abnormalities on MRI in the nasopharynx, deep face, or skull base should remain stable over this period or show further reduction in volume. Recurrent disease is seen as any increase in the bulk of abnormal signal from baseline imaging. Most recurrences, local or systemic, occur within the first 2 years after treatment (Glastonbury, 2007). Of patients with recurrences, 10% to 20% may be curable with additional treatment (Glastonbury, 2007). After 2 years without evidence of recurrence, the imaging interval is typically extended to every 6 to 12 months. Although not yet the mainstay of treatment, one meta-analysis demonstrated that FDG-PET/CT is more sensitive and specific than CT and MRI for the diagnosis of local residual or recurrent NPC (Liu et al., 2007).

### 5.1 MRI technique

The staging MRI protocol for NPC varies from center to center. In general, the images should cover the area from above the frontal sinuses to the thoracic inlet on axial studies and from the tip of the nose to the fourth ventricle on coronal sequences. At our institution, axial and coronal T1- and T2-weighted images, as well as a sagittal T1 or T2 fat saturation series covering the entire head and neck are obtained. A head and neck imaging coil is routinely used for both the 1.5T and 3.0T MRI scanners. The axial, coronal and sagittal T1 series are performed using a T1-FLAIR technique. Postgadolinium-enhanced axial images with fat saturation and coronal images without fat saturation are also acquired using a conventional spin echo T1 technique. Lau et al. found that the axial precontrast and postcontrast series were the most informative MRI sequences for evaluating primary tumour extension and achieved approximately 100% diagnostic accuracy in T-staging of NPC (Lau et al., 2004). We find that axial and coronal noncontrast T1-weighted images are
the best for providing detailed views of the local NP anatomy and surrounding structures, whereas postcontrast images allow for accurate assessment of perineural disease along major nerves, such as the maxillary and mandibular divisions of the trigeminal nerve, as well as the cavernous sinus (Yu et al., 2010).

5.2 Comparison of imaging methods

<table>
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<tr>
<th>MRI</th>
<th>CT</th>
<th>PET/CT</th>
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<tr>
<td><strong>Strengths</strong></td>
<td><strong>CT</strong></td>
<td><strong>PET/CT</strong></td>
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<tr>
<td>Superior to CT and PET/CT for assessing primary tumour invasion into surrounding soft tissue and bony structures, pharyngobasilar fascia invasion, invasion into the sinus of Morgagni, skull base invasion, intracranial invasion, as well as cavernous sinus extension and perineural disease (Liao et al., 2008; Ng et al., 2009 Sakata et al., 1999)</td>
<td>Relatively inexpensive</td>
<td>Superior to MRI and CT for assessing lymph node metastasis, especially cervical nodal metastases, and distant metastases, especially occult metastatic disease (i.e., N- and M-staging of NPC) (Comoretto et al., 2008; King &amp; Bhatia, 2010; King et al., 2008; Lin et al., 2008; Ng et al., 2009, Wang et al., 2007)</td>
</tr>
<tr>
<td>Superior to CT and PET/CT for evaluating retropharyngeal adenopathy</td>
<td>Rapid image acquisition time</td>
<td>Screens the entire patient for local recurrence, lymph node metastases and distant metastases during a single whole-body examination using a single injection of FDG</td>
</tr>
<tr>
<td>No exposure to ionizing radiation</td>
<td>Widely available</td>
<td>Significantly better sensitivity and specificity compared to CT and MRI for diagnosis of local residual or recurrent nasopharyngeal carcinoma (Liu et al., 2007)</td>
</tr>
<tr>
<td><strong>Weaknesses</strong></td>
<td><strong>CT</strong></td>
<td><strong>PET/CT</strong></td>
</tr>
<tr>
<td>Expensive</td>
<td>Less accurate than MRI and PET/CT for evaluating tumour invasion into surrounding tissues (Ng et al., 2009)</td>
<td>Accessibility issues (in some regions)</td>
</tr>
<tr>
<td>Long image acquisition times compared to CT</td>
<td>Exposure to ionizing radiation</td>
<td>Expensive</td>
</tr>
<tr>
<td>Less readily available compared to CT not always reliable in distinguishing between enhancing residual</td>
<td>Compared to MRI, PET/CT undermaps the</td>
<td>Exposure to ionizing radiation</td>
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6. Anatomy of the nasopharynx

The nasopharynx is at the superior and posterior aspect of the aerodigestive tract. The nasopharyngeal mucosa is lined with squamous epithelium and surrounded by a muscular and fascial sling consisting of the superior constrictor muscle and the buccopharyngeal fascia derived from the middle layer of the deep cervical fascia. The space has three walls and a roof. It opens anteriorly to the posterior nasal cavity via the posterior nasal choanae, and inferiorly into the oropharynx at the level of the hard palate. On imaging, the C1/2 junction is also an accepted marker between the nasopharynx and oropharynx (Dubrulle et al., 2007). The roof of the nasopharynx abuts the sphenoid sinus floor, and slopes posteroinferiorly along the clivus/basiocciput to the upper cervical vertebrae. Remains of adenoid tissue may persist into adulthood and exist as tags in the roof of the nasopharynx.

The pharyngobasilar fascia (PBF), a tough aponeurosis connecting the superior constrictor muscles to the skull base, is perhaps the most important structure of the nasopharynx. The tough fibers of the pharyngobasilar fascia create a framework that determines the configuration of the nasopharynx and the support by which the entire pharynx hangs from the skull base (Dillon et al., 1984). The parallel lateral walls of the pharyngobasilar fascia extend from the posterior margin of the medial pterygoid plate anteriorly to the occipital pharyngeal tuber and prevertebral muscles posteriorly. The foramen lacerum, which is...
within the confines of the pharyngobasilar fascia, is a fibrocartilaginous structure that forms part of the floor of the horizontal carotid canal and roof of the nasopharynx. It provides a route for nasopharyngeal tumors to access the cavernous sinus and intracranial cavity. The pharyngobasilar fascia is a fibrous structure and occasionally can be seen as a thin dark line on T2-weighted axial MR images, deep to the submucosal tissues of the nasopharynx (Figure 3).

Fig. 3. Axial T2 weighted image at the level of the nasopharynx. The arrow shows the pharyngobasilar fascia on the left side.

The paired Eustachian tubes, along with the medial fibers of the levator veli palatini muscle pass into the nasopharynx via the sinus of Morgagni, a posterolateral defect in the pharyngobasilar fascia. The opening of the Eustachian tube is anterior (on axial images) and inferior (on coronal images) to the torus tubarius, the distal cartilaginous end of the eustachian tube that forms a mucosal-lined structure projecting into the lumen of the nasopharynx from the superior aspect of the posterior lateral nasopharyngeal walls. The fossa of Rosenmüller, otherwise known as the lateral pharyngeal recess, is posterior (on axial images) and superior (on coronal images) to the torus tubarius. The fossa of Rosenmüller is the most common site of origin of NPC (Goh & Lim, 2009) (Figure 4). However, asymmetry of the lateral pharyngeal recesses is a common and normal incidental finding, and should not be mistaken as tumours.

Lateral to the nasopharynx is the parapharyngeal space (PPS), a fibrofatty space which separates the nasopharynx from the masticator space (Figure 4). Involvement of the parapharyngeal fat serves as an important marker of tumour infiltration used in staging. The posterolateral boundary of the nasopharynx consists of the carotid space (post-styloid parapharyngeal space), which is located posterior to the parapharyngeal space. Located posterior to the nasopharynx, between the nasopharyngeal mucosal space and the prevertebral muscles, is the retropharyngeal space, a potential space that contains the medial and lateral retropharyngeal lymph nodes. The lateral retropharyngeal nodes, also
known as the nodes of Röuviere, are the first nodes in the lymphatic drainage of the nasopharynx and, along with the cervical Level II nodes, are reported to be the most common site of nodal metastases (Wang et al., 2009). The lateral retropharyngeal lymph nodes can be identified on MRI from the skull base to the level of C3 (King & Bhatia, 2010). The medial retropharyngeal nodes do not form a discrete nodal chain, and thus, are less often visible on imaging.

![Fig. 4. Axial T1 image shows the torus tubarius (*). The solid arrow points to the opening of the Eustachian tube. Dashed arrow is the fossa of Rosenmuller. The dashed area shows the fat in the left prestyloid parapharyngeal space.](image)

Other important structures include the foramen rotundum and pterygoid (or Vidian) canal, which communicate with the pterygopalatine fossa and are potential routes of tumour spread.

### 7. Staging of NPC

After the initial diagnosis of NPC is made on history, physical examination, and biopsy, cross-sectional imaging is required for cancer staging. Currently, the 7th edition of the International Union Against Cancer (UICC) and American Joint Committee on Cancer (AJCC) TNM staging system is used, which was recently revised and released on 1 January 2010. As previously described, most cases of NPC originate in the lateral pharyngeal recess and spread submucosally with early infiltration into deeper neck spaces. NPC tends to have well-defined patterns of spread.
**Box 1.**

**TMN Staging of Nasopharyngeal Carcinoma**

**Overview**

**Primary Tumor (T)**

- **T1**: Tumor confined to the nasopharynx, or extends to oropharynx and/or nasal cavity without parapharyngeal extension
- **T2**: Tumor with parapharyngeal extension
- **T3**: Tumor involves bony structures of skull base and/or paranasal sinuses
- **T4**: Tumor with intracranial extension and/or involvement of cranial nerves, hypopharynx, orbit, or with extension to the infratemporal fossa/masticator space

*Parapharyngeal extension denotes posterolateral infiltration of tumor.*

**Regional Lymph Nodes (N)**

- **NX**: Regional lymph nodes cannot be assessed
- **N0**: No regional lymph node metastasis
- **N1**: Unilateral metastasis in lymph node(s), 5 cm or less in greatest dimension, above the supraclavicular fossa, and/or unilateral or bilateral, retropharyngeal lymph nodes, 6 cm or less, in greatest dimension
- **N2**: Bilateral metastasis in lymph node(s), 6 cm or less in greatest dimension, above the supraclavicular fossa
- **N3**: Metastasis in a lymph node(s) >6 cm in dimension and/or extension to supraclavicular fossa
- **N3a**: Greater than 6 cm in dimension
- **N3b**: Extension to the supraclavicular fossa

*Midline nodes are considered ipsilateral nodes.*

**Distant Metastasis (M)**

- **MX**: Distant metastasis cannot be assessed
- **M0**: No distant metastasis (no pathologic M0, use clinical M to complete stage group)
- **M1**: Distant metastasis

**AJCC Stage Groupings**

<table>
<thead>
<tr>
<th>Stage</th>
<th>Tis, N0, M0</th>
<th>Stage IVA</th>
<th>T4, N0, M0</th>
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<tr>
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<td>Stage IVB</td>
<td>T4, N1, M0</td>
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<td>Stage IVC</td>
<td>T4, N2, M0</td>
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<td>T2, N1, M0</td>
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<td>Any T, Any N, M1</td>
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<tr>
<td>Stage III</td>
<td>T3, N0, M0</td>
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<td>Any T, N2, M0</td>
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7.1 T1 NPC

Stage T1 refers to disease that is localized to the nasopharynx, as well as disease that has extended inferiorly into the oropharynx or anteriorly into the nasal cavity (Lee, 2004; Liu et al., 2008; Yu et al., 2010). Disease that is isolated to the NP is described as being superficial to the pharyngobasilar fascia (PBF). In this case, there is no evidence of tumour breaching or crossing the PBF barrier, which occasionally can be identified on T2-weighted studies as a thin dark curvilinear line running from the medial pterygoid plate laterally and following the contour of the longus musculature along the back wall of the NP (Figure 3). The tumour appears as a hypointense to isointense mass on T1-weighted imaging that enhances to a lesser degree than normal mucosa (Chin et al., 2003).

Oropharyngeal extension is readily noted on coronal or sagittal MR imaging as tumour that has extended inferiorly past the plane of palate (Figure 5). On axial sections, the oropharynx is considered involved when tumour is seen inferior to the C1/C2 junction.

Fig. 5. Axial image of a patient with NPC that shows disease that has extended down to the level of the right oropharynx.

In addition, stage T1 disease comprises anterior extension past the plane of the posterior choana into the nasal cavity (Figure 6). From the nasal cavity, NPC can invade through the sphenopalatine foramen into the pterygopalatine fossa (stage T3 disease), resulting in obliteration of the normal fat content in this fossa (Figure 7). Direct extension from the nasal cavity is the most common route of NPC invasion into the pterygopalatine fossa, followed
by extension from the ethmoid and/or sphenoid sinuses. The significance of pterygopalatine fossa involvement is described later (T3 NPC).

Fig. 6. NPC which has extended beyond the plane of the posterior choana (dashed line) into the nasal cavity.

Fig. 7. Axial T1 weighted image shows abnormal soft tissue filling the right and left pterygopalatine fossae (*).
7.2 T2 NPC

Disease that has spread beyond the PBF and infiltrated posterolaterally into the parapharyngeal space (PPS) is considered stage T2. This can lead to compression of the Eustachian tube, resulting in middle ear and mastoid effusion (King & Bhatia, 2010). PPS invasion is associated with worse disease control and survival rates, as well as increased rate of distant metastases compared to stage T1 (Cheng et al., 2005; Chua et al., 1996; Ho et al., 2008; Teo et al., 1996). On imaging, key components of the PPS include the tensor veli palatini and levator palatine muscles. PPS involvement is recognized as a disruption or attenuation of the PBF, infiltration of the tensor veli palatini, or breach of the intrapharyngeal portion of the levator veli palatini muscle.

Infiltration of the fatty component of the PPS is visible on T1-weighted imaging as an intermediate signal mass invading into the hyperintense fat of the PPS (Figure 8). As the disease progresses, the full thickness of the intrapharyngeal portion of the levator veli muscle and the tensor veli palatini muscle (appearing as a thin strip of muscle just lateral to the levator veli palatini) may also become involved, followed by infiltration of the normally pristine fat of the prestyloid PPS, which is located deep to the tensor muscle. With further posterolateral spread, the poststyloid PPS structures, such as the carotid sheath, also become vulnerable to disease (Figure 9).

Fig. 8. Axial T2 weighted images shows a carcinoma that has infiltrated lateral to the tensor and levator veli palatini. Solid arrow shows normal tensor veli palatini on the right. Dashed arrow shows the abnormal left levator muscle which is being displaced and the presence of tumor lateral to it.
Stage T3 disease is characterized by involvement of the paranasal sinuses and/or the central skull-base structures. Skull base invasion is seen in up to 60% of NPC patients at diagnosis (King et al., 1999; Roh et al., 2004), with the most common sites of involvement being the clivus, pterygoid bones, body of the sphenoid bone, and apices of the petrous temporal bones (King & Bhatia, 2010). In addition, the number of sites involved within the skull base may have prognostic significance (Lu et al., 2004). Assessment of the skull base should focus on five key regions: clivus, right pterygoid base, left pterygoid base, right petrous apex, and left petrous apex (King & Bhatia, 2010). On axial T1-weighted MR sequences, bony skull base disease is detected as a loss of the hyperintense signal, which is characteristic of normal fatty yellow bone marrow. It is replaced with an intermediate signal indicative of either bony reaction or actual invasion. NPC commonly invades posteriorly into the clivus (Figure 10). Although normal clival marrow may appear heterogeneous on T1-weighted images, it should still appear more intense than the pons (Goh & Lim, 2009). In addition, up to 25% of patients have tumour spread superiorly into the floor of the sphenoid sinus, often extending up into the sphenoid sinus cavity (Chong & Fan, 1993; Goh & Lim, 2009) (Figure 11). The body of the sphenoid is more difficult to assess and may be best studied on coronal images (King & Bhatia, 2010). The sphenoid wings and upper cervical spine should also be assessed. CT scans may also have a role in detecting the presence of cortical erosion and bony sclerosis, which reflects reactive changes due to tumour invasion. It is important, however, to note that sclerosis is a non-specific sign that may arise from adjacent sinus disease, particularly in the sinus margins and the pterygoids.
Fig. 10. 10A is a sagittal T1 image that shows NPC invasion into the clivus. There is a loss of the inferior cortical margin of the clivus. The clivus is also of uniform lower signal that reflects sclerotic change. 10B is a coronal T1 image in another patient that shows NPC infiltration into the left aspect of the clivus (arrow).

Fig. 11. Coronal T2 (left) and post gadolinium enhanced T1 (right) weighted images show superior extension of NPC through the floor of the sphenoid sinus (arrows).
Next, the skull base foramina and fissures – which include the foramen rotundum (V2 nerve), the vidian canal (vidian nerve), the foramen ovale (V3 nerve), and foramen lacerum – should be examined. Nerve involvement signifies stage T4 disease (see section below). The foramen ovale and lacerum are common routes of tumour extension into the intracranial cavity (King & Bhatia, 2010). While the skull base foramina present an unobstructed route for tumor spread, direct invasion of the bone bordering these foramina is also a common finding. The skull base foramina are best assessed on coronal images. Less common findings include inferior spread of tumor to involve the hypoglossal nerve canal (XII nerve) and jugular foramen (IX-XI nerves) (King & Bhatia, 2010).

It is important to differentiate tumour spread into the paranasal sinuses – which include the ethmoid sinuses, sphenoid sinuses, and maxillary sinuses – from benign mucosal inflammatory changes, which are common in patients with NPC. On MRI, mucosal thickening can be distinguished from NPC by a greater uniform T2-weighted signal and a marked degree of contrast enhancement. With tumor invasion into the ethmoid sinuses, which usually occurs via direct spread from the sphenoid sinus or nasal cavity, there is a reduced chance of shielding the optic nerve from the radiation dose in RT. Involvement of the sphenoid sinus, which is separated from the nasopharynx by only a thin plate of bone, is a common finding (Figure 11). In contrast, anteroinferior tumour invasion into the maxillary sinuses is a rare event, except in late disease where there is usually extensive invasion throughout the nasal cavity, other paranasal sinuses, skull base, and brain (King et al., 1999). Sinus involvement is seen on imaging as a loss of contiguity of the sinus walls.

### 7.4 T4 NPC

T4 disease is denoted by intracranial extension and/or involvement of the cranial nerves, hypopharynx, orbit, or masticator space. Unchecked, NPC can extend superiorly into the cavernous sinus and dura (Figure 12), while direct invasion of the brain is rare at diagnosis (King & Bhatia, 2010). There are multiple routes into the cavernous sinus, including extension from tumour surrounding the horizontal portion of the internal carotid artery or cranial nerve V3, as well as extension from the orbital fissures or the skull base in the region of the foramen ovale or sphenoid sinus (King & Bhatia, 2010). Within the cavernous sinus, cranial nerves III, IV, V1, V2, and VI are vulnerable to tumour invasion. Their involvement is manifested clinically through an abnormal neurological examination, as well as by signs and symptoms of various cranial nerve palsies, including extraocular muscle dysfunction, facial pain, paresthesia and numbness. True perineural spread, however, is uncommon in the pre-treatment setting (King & Bhatia, 2010). Prognostically, cranial nerve involvement has been shown to be associated with a higher rate of distant metastases and decreased survival (Yu et al., 2010). Radiologically, post-contrast T1-weighted MR sequences with fat saturation are used to assess for cranial nerve involvement, especially for V2 along foramen rotundum (Figure 13) and V3 in the foramen ovale (Figure 14). Contrast-enhanced MRI shows perineural disease extension and cavernous sinus involvement as asymmetric nodular thickening and abnormal enhancement. Skip lesions may also be noted. A late sequelae is expansion of the bony canals in which these nerves travel.
Imaging of Nasopharyngeal Carcinoma

Fig. 12. Coronal postcontrast study shows a large NPC that has extended laterally into the right masticator space as well as superiorly to invade the cavernous sinus, sphenoid sinus and cause mass effect upon the right temporal lobe.

Fig. 13. Axial image shows asymmetric thickening and enhancement along the left V2 nerve (arrows).

Orbital invasion, usually arising from the pterygopalatine fossa via the inferior orbital fissure or directly from the cavernous sinus, denotes the most extensive form of disease (King & Bhatia, 2010) (Figure 15). From the PPS, pterygoid base, or pterygomaxillary fissure, NPC can extend laterally to involve masticator space, which includes the medial and lateral pterygoid muscles, temporalis muscle, infratemporal fat, as well as the mandibular division (V3) of the trigeminal nerve (Figure 14). Disease involving the masticator space can give rise to trismus and weakness in mastication. In addition, the V3 nerve is vulnerable to tumour infiltration within the masticator space. Both antegrade and retrograde perineural spread along V3 is possible. From the masticator space, NPC can also extend superiorly through the floor of the middle cranial fossa and foramen ovale to gain access to the intracranial cavity and the cavernous sinus.
Fig. 14. Coronal image shows NPC that has extended laterally into the left masticator space and subsequent perineural tumor tracking superiorly (dashed arrows) along V3 into the left cavernous sinus (*). Solid arrow shows the normal contralateral V3 nerve.

Fig. 15. Axial T2 weighted scan shows intermediate signal tumor that has extended through the superior orbital fissure into the left orbit.

The hypopharynx, which is the most inferior site included in the TMN staging classification, is very rarely involved at diagnosis due to the tendency of NPC to extend superiorly rather than inferiorly (King & Bhatia, 2010).

7.5 Nodal stage

The NPC N classification system differs from other head and neck mucosal malignancies (see Box 1). Up to 60–90% of NPC patients will have nodal metastases at presentation (Glastonbury, 2007; Goh & Lim, 2009), suggesting that only 10-40% of cases present
without positive nodal disease (N0). Positive neck nodal disease in NPC is associated with an increased risk of local recurrence and distant metastases (Goh & Lim, 2009). Unlike other head and neck squamous cell carcinomas, nodal disease in NPC is more frequently bilateral.

NPC generally follows a very orderly pathway of nodal spread, beginning with the (lateral) retropharyngeal lymph nodes (RPN) (Figure 16) – located medial to the carotid artery – before involving nodal groups along the internal jugular chain (level II to IV), spinal accessory chain (Va and Vb), as well as supraclavicular nodes (Glastonbury, 2007; King et al., 2004). Level IIa or b nodes, located posterior to the jugular vein in the upper neck, are the most common site for non-retropharyngeal node involvement (Mao et al., 2008) (Figure 17).

Fig. 16. Cystic right retropharyngeal lymph node.

Fig. 17. Right sided level II adenopathy (*).
Nodal disease in the submandibular and parotid/periparotid regions is a rare occurrence (Chong & Fan, 2000; King & Bhatia, 2010), but should be evaluated for radiotherapy planning purposes. Although the RPN are generally considered the first echelon of metastatic spread, studies have shown that this is not true in all cases (Liu et al., 2006; Mao et al., 2008; Ng et al., 2004; Wang et al., 2009). While King et al. found that the RPN were bypassed in only 6% of cases to preferentially affect the level II distribution, Ng et al. found that 17 of 89 cases of NPC bypassed the RPN (King et al., 2000; Ng et al., 2004). In addition, Ng et al. also reported skip metastases in the lower neck lymph nodes and the supraclavicular fossa in 7.9% of cases, and distant metastases to thoracic and abdominal nodes in 3-5% of cases. After radiotherapy, level I nodes may also become involved (Ahuja et al., 1999).

In the current N classification system, N1 disease consists of the presence of unilateral metastasis in cervical lymph node(s) that are 6 cm or less in their greatest dimension, and/or unilateral or bilateral RPN metastasis that are 6 cm or less in their greatest dimension. Nodes greater than 3 cm in size are generally considered “nodal masses” and are indicative of confluent nodes. N2 disease is defined by bilateral metastasis in the cervical lymph node(s) that are again 6 cm or less in greatest dimension. Unlike other carcinomas in the neck, N2 is not further divided into substages. In both N1 and N2 stages, disease is restricted to above the supraclavicular fossa. Once supraclavicular fossa nodal involvement is noted, the disease is upstaged to N3b. Supraclavicular nodes include all lymph nodes seen on the same axial cross-sectional slice as a portion of the clavicle, and include Level IV and Vb nodes. Stage N3a disease is defined by matted nodes greater than 6 cm in their greatest dimension, although this is a rare finding (King & Bhatia, 2010) (Figure 18).

Fig. 18. Massive conglomerate right sided neck adenopathy measuring over 6cm in dimension -N3a disease.

While distinguishing between the primary tumour mass and adjacent RPN is best assessed with MRI, cervical lymph nodes can be evaluated accurately with both MRI and CT (King et al., 2004). T2-weighted imaging with fat saturation shows nodes as bright structures in the posterior cervical fat (Figure 17). The higher resolution of CT facilitates the visualization of neck adenopathy, as well as nodal necrosis and extracapsular extension. The latter manifests
as a loss or irregularity of the nodal margins, and/or streakiness of the adjacent fat (Yu et al., 2010). Patients with nodes showing necrosis and extranodal spread have a very poor prognosis with a 50% decreased 5-year survival rate (Som et al., 1987).

There are several features on imaging that are suggestive of metastatic nodal disease, including a large size (although there is no accepted size criterion, generally >1.5 cm for levels I and II, >1 cm for levels IV-VII, and >5 mm for RPN using the shortest transaxial diameter is considered suspicious) (Goh & Lim, 2009; King & Bhatia, 2010), a group of 3 or more nodes borderline in size, rounded nodes with loss of the fatty hilum, and necrosis (King & Bhatia, 2010). If identified, necrosis is considered 100% specific. However, due to resolution restrictions, necrosis can only be reliability identified in tumour foci greater than 3 mm, of which approximately one-third reportedly have nodal necrosis (Goh & Lim, 2009; Som & Brandwein, 2003; Yousem et al., 1992). Necrosis or cystic change is hypointense on T1-weighted images with rim enhancement with contrast, and hyperintense on T2-weighted images (Figure 16). In CT images, necrosis is seen as a focal area of hypoattenuation with or without rim enhancement (Figure 19).

Fig. 19. Bilateral level II adenopathy. Focal areas of lower attenuation on CT is compatible with the presence of necrotic change.

7.6 Distant metastasis (M)

The M stage of NPC is similar to that of other malignancies, whereby M0 signifies the absence of distant metastasis and M1 refers to the presence of such disease. NPC has the highest incidence of distant metastasis among head and neck cancers, with a rate as high as 11% at diagnosis (Kumar et al., 2004; Teo et al., 1996). The likelihood of metastasis increases with increasing T and N stage.
The most commonly affected regions are the bone, lung and liver (Chiesa & De Paoli, 2001) (Figure 20). Thus, bones and lung apices should be evaluated for tumour involvement in head and neck MRI studies, especially in patients with risk factors such as metastatic cervical nodes which extend to the supraclavicular fossa (stage N3b).

The presence of M1 disease is associated with shorter survival rates (Teo et al., 1996) and significantly alters patient management, as such patients are generally considered to be incurable. Median survival is under four months and approximately 90% of patients with distant metastases pass away within a year (Goh & Lim, 2009; Khor et al., 1978). The goal of treatment in such instances will be palliation with therapy being applied for locoregional symptom control.

The exact method for the evaluation of distant metastasis will vary from institution to institution. Imaging options include bone scintigraphy, chest x-ray, CT of the thorax, abdomen and pelvis, and PET/CT. Studies have shown fluorodeoxyglucose PET/CT imaging to have a higher sensitivity and specificity in detecting distant metastases (Chen et al., 2006; Chua et al., 2009; Comoretto et al., 2008; King et al., 2008; Lin et al., 2008; Liu et al., 2007; Ng et al., 2009; Wang et al., 2007; Yu et al., 2010).
8. Tumour volume

Tumour volume has also been reported to have significant prognostic relevance outside of the TNM classification system (Wei & Sham, 2005). There is an estimated 1% increase risk of local control failure with every 1 cm³ increase in tumour volume. The increased risk is attributed to factors such as increase number of tumour clonogens, tumor hypoxia and relative radioresistance, etc (Bentzen et al., 1991; Johnston et al., 1995; Lartigau et al., 1993; Sze et al., 2004). However, issues with standardized methods of volume measurement, the intra- and interobserver reliability due to operator-dependent tracing, as well as the technical challenges associated with implementing this in the clinical setting – including the time it takes to perform this tedious task – have prevented it from being routinely used in daily practice.

9. Summary

Nasopharyngeal carcinoma is a relatively rare neoplasm with a characteristic geographic and ethnic distribution. It most commonly arises in the lateral pharyngeal recess, and has a tendency to invade widely and metastasize. Cervical lymphadenopathy is very common at presentation and is a common presenting complaint.

Diagnosis of NPC can be made on endoscopically-guided biopsy, but effective treatment of NPC requires an accurate mapping of tumor volume and extent with imaging. Imaging allows for evaluation of tumour extent, including submucosal, osseous, and intracranial tumour spread which cannot be assessed clinically or endoscopically. Staging of NPC is based on the new 2010 AJCC guidelines. MRI is the best tool for assessing tumour extent under the current staging system, while high-resolution CT has value for assessing cortical bone erosion and PET/CT is emerging as the most accurate modality for assessing distant metastases and recurrence.

A thorough understanding of the anatomy of the nasopharynx and surrounding structures, as well as the natural history of the disease and patterns of spread, allows for accurate tumour mapping and treatment planning.

10. References


Chen YK, Su CT, Ding HJ et al. (2006). Clinical usefulness of fused PET/CT compared with PET alone or CT alone in nasopharyngeal carcinoma patients. Anticancer Res. 26(2B), 1471–1477.


This book is a comprehensive treatise of the potential risk factors associated with NPC development, the tools employed in the diagnosis and detection of NPC, the concepts behind NPC patients who develop neuroendocrine abnormalities and ear-related complications after radiotherapy and chemotherapy, the molecular mechanisms leading to NPC carcinogenesis, and the potential therapeutic molecular targets for NPC.

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