Management of malignant tumors of the anterior skull base: experience with 76 patients

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As the management of anterior cranial fossa malignancies has undergone significant evolution, decreases in morbidity and mortality rates have occurred. In this article, the authors discuss the clinical presentation, neuroimaging findings, and management options for common anterior skull base malignancies. Also discussed are surgery-related indications and principles.

KEY WORDS  •  anterior skull base  •  transbasal  •  nasopharyngeal carcinoma  •  esthesioneuroblastoma  •  transfacial

In 1941 Dandy described anterior craniofacial resection of orbital tumors.20 Prior to this, Frazier21 championed removing orbital rims to gain improved surgical access. In 1943 Ray and McLean59 combined a transcranial and transfacial approach for resection of a retinoblastoma. The first en bloc resection of an extensive paranasal sinus malignancy was initially reported by Smith, et al.,71 in 1954 and subsequently by Ketcham, et al.,38 and Van Buren, et al.79 Although complete tumor resection had considerable benefits, the toll was an 80% morbidity rate and a 7% mortality rate.38,69 The major cause of morbidity was intracranial contamination due to sinonasal tract communication.64 This complication was reduced when reconstructive techniques involving pericranial grafts were introduced in 1979 by Schramm, et al.,65 and modified later by Johns, et al.37 With improvements in surgical techniques, the morbidity rate decreased to less than 40% and mortality to between 0 to 2%.45,46

In 1960 Tessier, et al.,76 pioneered the transbasal approach for the reduction of craniofacial abnormalities. Derome, alone,21 with Tessier,22 and with others23 used this approach for sphenoidvenous tumor removal beginning in the 1970s. Modifications of this approach have led to increased anterior cranial fossa exposure requiring minimal brain retraction.

Anterior transfacial approaches were described as early as 1829 by Lizars. Modifications were described by Fergusson in 1842.9 In the 1920s, Portmann and Retrouvey reported the midfacial degloving procedure.7 Casson and associates10 further modified this technique, which was then popularized by Conley and Price16 and Price alone.55 A subcranial approach obviating the need for the transfacial approach was initially described by Raveh, et al.,58 in 1978 for the management of craniofacial fractures and subsequently performed for the management of anterior skull base tumors.

The management of malignant tumors of the anterior skull base can be technically demanding and has been associated with high rates of morbidity and mortality. Advances in neuroimaging leading to earlier diagnosis, a better understanding of the anatomy, and collaboration among experienced surgeons has led to significant progress in the management of these lesions.

MANAGEMENT OF ANTERIOR SKULL BASE MALIGANCIES

Presentation and Neuroimaging Features

Anterior fossa tumors can manifest as a number of different symptoms, including nasal obstruction and congestion, epistaxis, rhinorrhea, hyposmia or anosmia, headache, seizure, and psychological changes. Paresthesias in the distribution of the V1 or V2 divisions of the trigeminal nerve may occur. Invasion into the sella turcica can cause loss of endocrine function, or invasive pituitary tu-
mors can be responsible for over production of corticosteroids, growth hormone, or prolactin. Visual loss can be uni- or bilateral. Associated ocular symptoms include diplopia, orbital pain, exophthalmos, and retroorbital headaches. Many of these tumors can be demonstrated on nasopharyngeal examination.

Diverse tumors can involve the anterior cranial fossa. Primary benign tumors with rare malignant potential include meningiomas and pituitary adenomas. Primary malignant tumors include chordomas, chondrosarcomas, osteogenic sarcomas, and invasive pituitary tumors. Malignant tumors with secondary involvement include carcinoma of the paranasal sinuses. Others include lymphoma arising from the adenoids, minor salivary gland neoplasms, ethmoid carcinoma, paragangliomas, olfactory neuroblastomas, mucoepidermoid carcinomas, and osteosarcomas. Histologically benign tumors include angiobromas and inverted papillomas. Metastases from renal, breast, and lung carcinoma can also involve the anterior cranial fossa. Teratocarcinomas are rare in this region (Fig. 1). Lastly, paranasal sinus infections can extend into the anterior skull base.

Computerized tomography and MR imaging provide complementary information regarding skull base lesions. Whereas CT scanning is better at defining osseous anatomy and calcifications, MR imaging is superior at demonstrating the relationship of skull base masses to soft-tissue structures such as cranial nerves, the CA, the jugular vein, and the brain. In the case of parasellar masses, MR imaging is better in excluding CA aneurysms.

Nasopharyngeal Carcinoma. Nasopharyngeal carcinoma arises from the surface epithelium of the nasopharynx and commonly metastasizes to the lymph nodes, with some preference for those at the apex of the posterior cervical triangle.18 As many as 82% of the tumors arise in the lateral wall of the nasopharynx in Rosenmüller fossa (pharyngeal recess), 12% in the midline, and 6% in normal-appearing mucosa.70

Nasopharyngeal carcinoma has a bimodal age distribu-
tion with peaks in the second and sixth decades of life. It occurs predominantly in males. It is divided into two distinct histological entities: squamous cell carcinoma and UCNT. The WHO classifies nasopharyngeal carcinoma based on the degree of differentiation:47,70,81 Type I is a keratinizing squamous cell carcinoma similar to that found in the rest of the upper digestive tract; Type II is a nonkeratinizing epidermoid carcinoma; and Type III is UCNT. Numerous staging schemes are present for maxillary and ethmoid neoplasms; however, only the University of Florida staging system can be applied to all paranasal neoplasms.50,52 In this classification system, T1 tumors are limited to a single site of origin such as a nasal wall or ethmoids; T2 tumors extend to an adjacent sinus, orbit, palate, nasopharynx, or pterygomaxillary sites; and T3 tumors extend to or through the skull base. Resection of T1 tumors is associated with the least morbidity and mortality.

The UCNT differs from other head and neck squamous cell carcinomas by its histological features, epidemiological characteristics, and its relation with the Epstein–Barr virus. Of nasopharyngeal carcinomas, it is the more prevalent. It is endemic in Southeast Asia, parts of the Mediterranean basin, North Africa, parts of the Caribbean and Alaska. A characteristic feature of the undifferentiated tumor is massive lymphoid T-cell infiltration.

The UCNT is associated with the Epstein–Barr virus.47,75,78,80 Deletion of tumor suppressor genes such as p16 or the activation of oncogenes such as bcl-2 may predispose individuals to the development of WHO Types II and III carcinoma.47,48 Dietary factors such as salted fish high in volatile nitrosamines have also been implicated in UCNT.82

Squamous cell carcinoma (WHO Type I) is more common in the Western population. The use of tobacco and alcohol may play a role in the development of this well-differentiated tumor.13 It is rarely associated with the Epstein–Barr virus. Mutations of the p53 gene commonly occur.16

Patients with nasopharyngeal carcinoma may present with variable nonspecific symptoms, the most common of which is painless posterior cervical adenopathy. Lateral pharyngeal node involvement may result in pain with ipsilateral neck rotation or ear pain. Unilateral conductive hearing loss resulting from serous effusions due to poor eustachian tube function commonly occurs. Nasal obstruction occurs late in the course of the disease and profuse epistaxis is rare. Dysphonia, difficulty swallowing, and diplopia may also occur. Physical examination may reveal maxillary nerve dysfunction. Extension of the tumor into the lateral pharynx results in ninth and 10th cranial nerve palsies. Proptosis and trismus are rarely associated with these malignancies.

Nasopharyngeal carcinoma is associated with bone destruction demonstrated on CT scanning. Intense homogeneous enhancement of the lesion is seen following contrast administration. The lesion can invade the middle fossa either by erosion through the greater wing of the sphenoid or extension through the foramen ovale. In advanced cases, the cancer can reach the clivus and violate areas of the posterior fossa. Adenoid cystic carcinomas are also associated with bone destruction.
Anterior skull base malignancies

MR imaging may reveal perineural spread of the malignancy particularly along the V2 and V3 distribution.

The treatment of choice for nasopharyngeal carcinoma is chemotherapy followed by radiotherapy, although some authors have advocated surgery followed by radiotherapy. The histological type clearly affects outcome. Patients with squamous cell carcinoma fare worse than those with WHO Type II or III carcinoma in terms of local disease control and overall survival. The Type II and III carcinomas are more radio- and chemosensitive compared with the keratinizing squamous cell carcinoma. Despite improvements in management of nasopharyngeal carcinomas, the rate of cancer-free survival at 5 years is less than 50% for all stages. It is 25 to 30% for advanced stages of disease.

Although early results suggest that chemotherapy may be beneficial, the data are limited. Providing concomitant radio- and chemotherapy with cisplatin, Choi, et al. reported 5-year survival rates in 50% of their patients with advanced disease. Induction chemoradiotherapy with 5-fluouracil and cisplatin may lead to improved survival rates. Intraarterial chemotherapy has recently been applied in cases of paranasal sinus cancers supplied by the internal maxillary artery. Selective delivery of agents such as cisplatin by this technique increases tumor–drug concentration.

Olfactory Neuroblastoma (Esthesioneuroblastoma). Olfactory neuroblastomas, also known as esthesioneuroblastomas, are rare tumors arising from olfactory mucosa of the cribiform plate, superior turbinate, and the upper third of the nasal septum. They present most commonly with intractable nasal blockage or epistaxis, less commonly with local headache, rhinorrhea, visual disturbances, or anosmia, and rarely with proptosis, cheek swelling, or a neck mass. Modified Kadish system tumor staging provides the following classifications: Stage A, confined to nasal cavity; B, involving area around paranasal sinus; C, extension in the orbit, skull base, or cranium; and D, distant metastases. Cervical metastases occur in fewer than 10% of Stage A and B cases but are present in 44% of Stage C cases.

Evaluation should include CT evaluation to evaluate osseous erosion of the skull base as well as MR imaging to differentiate tumor from sinus obstruction in the paranasal sinuses (Figs. 2 and 3). Gadolinium enhancement is used to evaluate tumor erosion through the cribiform plate and metastases to cervical lymph nodes. The differential diagnosis for olfactory neuroblastoma includes melanoma, lymphoma, extramedullary plasmacytoma, rhabdomyosarcoma, and sinonasal undifferentiated carcinoma.

At present there is no proven optimal therapy for esthesioneuroblastoma, but a consensus does seem to be emerging in the literature. Lesions confined to the nasal cavity, free from the cribiform plate, and with tumor-free margins can be treated by surgery alone via an extracranial approach. Anterior craniofacial surgery is performed in the majority of cases involving the cribiform plate. The extracranial dissection can be performed transasally, endoscopically, or through a lateral rhinotomy. Intracranial surgery can be approached through a bifrontal craniotomy or through large frontal sinuses, taking advantage of a frontal sinus template prepared prior to surgery.

Radiotherapy is recommended for low-grade tumors without clean surgical margins and all high-grade tumors. Most radiotherapists recommend administering a 50 to 60-Gy dose of radiation postoperatively. At some institutions a 50-Gy dose is administered preoperatively to shrink large irregular tumors. Formulating a treatment plan for radiotherapy is difficult because of the irregular shape of the tumor and the adjacent radiosensitive optic nerves, pituitary gland, and frontal lobes.

The most commonly used chemotherapy regimen includes cyclophosphamide and vincristine. Electron microscopy shows neurosecretory granules, and immunohistochemistry is positive for S100 protein and neuron-specific enolase and negative for epithelial and lymphoma markers. The absence of rosettes, increased mitotic activity, necrosis, and nuclear pleomorphism in olfactory neuroblastoma correlate with a poor prognosis.

Rhabdomyosarcoma. Rhabdomyosarcomas of the head and neck were first reported in 1854 by Weber and not histologically classified until 1947. These tumors occur predominantly in the pediatric population. Children younger than age 5 years harbor the aggressive embryonal variant and older patients the pleomorphic variant. Approximately 40% of patients have head and neck involvement, which can be orbital, nonparameningeal/nonorbital, and parameningeal.

Magnetic resonance imaging reveals a soft-tissue mass in the sinonasal region, infraorbital fossa, or nasopharynx with significant contrast enhancement. Computerized tomography scanning commonly demonstrates bone destruction. Five-year survival rates of 94% in cases of orbital tumors and 50% in cases involving other head and neck regions were reported in 1983.

In recent years, improved outcomes have been associated with early provision of chemoradiation therapy and resection. Radiotherapy in 150- to 200-cGy fractions is commonly undertaken for a total dose of 4140 to 6500 cGy. Radiotherapy is primarily responsible for the neuroendocrine, dental, and thyroid sequelae. It partially contributes to visual loss and facial asymmetry. Investigators in the Intergroup Rhabdomyosarcoma Studies II and III...
have demonstrated that at least 77% of children develop one problem within 7 years following their treatment regimen. Long-term follow-up data in these patients are needed because the late effects of radiotherapy can occur up to 10 years posttreatment.

Recurrences when resection is not performed are common. Rhabdomyosarcomas have a propensity to metastasize prior to operative intervention. Pulmonary metastases are associated with the worst prognosis.

Chordomas and Chondrosarcomas. Chordomas were described as early as 1856 by Lusckha. They are malignant neoplasms arising from remnants of the embryonic notochord. Chordomas and chondrosarcomas, which account for fewer than 1% of intracranial tumors, are pathologically distinct. These tumors have a propensity to occur in the third to fifth decades of life. Chordomas and chondrosarcomas invade locally (Fig. 4). Although relatively rare, metastases of intracranial chordomas to the lymph nodes, lungs, liver, bone, skin, and muscles have been reported. The conventional chordoma has vacuolated mucus containing physaliphorous cells. The chondroid type has cartilaginous tissue and is associated with a better prognosis. Some authors have argued these are immunohistochemically low-grade chondrosarcomas.

Chondrosarcomas are classified as classic, mesenchymal, or dedifferentiated. The classic variant has large multinucleated cells with abundant cartilage. This group is further divided into Grades I to III based on mitoses, nuclear size, and extent of cartilage matrix. The mesenchymal variant has islands of undifferentiated mesenchymal cells and cartilage. These are more aggressive tumors. They commonly occur at the petrosphenocilal junction. The dedifferentiated variant is similar to anaplastic sarcoma. Immunohistochemical markers can differentiate chordomas and chondrosarcomas. The former do not stain for epithelial markers or oncotic antigens.

The most common presenting symptoms include orbitofrontal headaches, visual disturbances, and ophthalmoplegia. Trigeminal, facial, and vestibulocochlear neuropathies may also be present because of extension of the tumor into the cerebellopontine angle and internal auditory meatus. Computerized tomography scanning reveals bone destruction without sclerosis. Magnetic resonance imaging reveals hypointense or isointense lesions on T1-weighted sequences that enhance brightly with gadolinium. Cystic areas of hemorrhage or mucoid material may be present. These are bright on T2-weighted sequences.

In patients with chondrosarcomas who undergo surgery and proton-beam radiotherapy 5- and 10-year survival rates of 99 and 98%, respectively, have been reported. Chordomas are associated with a less favorable prognosis, their 5- and 10-year survival rates being 51 and 35%, respectively. The five-year recurrence-free survival rate following gross-total resection and proton-beam radiotherapy is 100% for chondrosarcomas and 51.6% for chordomas.

Chordomas and chondrosarcomas are resistant to conventional doses of radiotherapy in the range of 50 to 70 Gy. They are radiosensitive to doses as high as 80 Gy. Considerable injury to adjacent critical neurovascular structures and radiation-induced necrosis are the major morbidities. Proton-beam radiotherapy with dose ranges of 50- to 75-cobalt Gy equivalents, has been administered as an adjunctive therapy to microsurgical resection to improve outcomes. This treatment modality has a 34% complication rate. Endocrine, auditory, seizures, and radionecrosis complications occur. No chemotherapy regimen has proven consistently effective. Chemotherapy is
used as a last resort in patients in whom both surgery and radiotherapy have failed.

The consensus emerging from the literature is that resection followed by proton-beam radiotherapy for residual tumor is the most effective therapeutic paradigm. Total or near-total resection of these tumors by experienced surgeons does not increase the risk of postoperative cranial nerve deficits.

Malignant Meningiomas. Meningiomas of the anterior cranial fossa skull can be typical benign tumors, atypical tumors prone to recurrence, or rare lesions that are overly malignant or anaplastic. Three percent of all intracranial meningiomas secondarily involve the sinonasal tract. Tumor extension can also lead to involvement of the orbit and parapharyngeal space. Predisposing factors for the development of meningiomas include female sex, previous radiotherapy, and neurofibromatosis Type 2. Atypical meningiomas do not have frank anaplasia. They demonstrate hypercellularity, patternless growth, focal necrosis, high nucleus/cytoplasm ratios, coarse chromatin, and prominent nucleoli. Numerous tumor cells undergoing mitosis are also commonly seen. Increased mitotic activity in this group of tumors is defined as four or more mitoses per 10 hpf. Aneuploidy and high proliferation index based on bromodeoxyuridine studies correlate with a higher rate of local recurrence.

Malignant meningiomas, by definition, have unequivocal anaplasia and/or invasion of brain parenchyma. Neuroimaging demonstrates an inordinate amount of edema and lack of a smooth contoured tumor–brain interface. Cytologically, the malignant areas range from a meningothelial appearance to a vaguely epithelial or sarcomatous appearance. The meningiomatous nature of the anaplastic tumor may be difficult to appreciate initially. Twenty or more mitoses per 10 hpf are present. The most common sites of metastasis are liver, lungs, pleura, and lymph nodes. Increased labeling with bromodeoxyuridine and monoclonal antibody Ki-67 occur in cases of aggressive tumors. Meningiomas are positive for epithelial membrane antigen and vimentin, and negative for anti–Leu 7 and glial fibrillary acidic protein. In patients with malignant meningiomas median survival is less than 2 years.

The three common histological patterns of typical meningiomas are syncytial, fibroblastic, and transitional. The syncytial growth pattern is the most common. Histological patterns have no prognostic significance. The most common chromosomal abnormality is loss of chromosome 22 q 12.3-q ter. In meningiomas with an aggressive or invasive course and frequent recurrences deletion of the short arm of chromosome 1 is also present. Curative therapy for meningiomas is complete resection. Radiotherapy is reserved for application in patients in whom incomplete tumor resections have been performed because of the risk of injury to critical neurovascular structures; in those with malignant meningiomas; or in rare cases in which the tumor may be judged inoperable. A large number of nonmalignant meningiomas have hormonal receptors. Pharmacotherapy aimed at these hormonal receptors includes bromocriptine, mifepristone, and RU486. Although there was initial enthusiasm for antiprogesterones, its administration in patients with unresectable meningiomas has not been encouraging.

Trapidil has known antagonistic action on platelet-derived growth factor and has been shown to exhibit some inhibition of meningioma cell proliferation.

SURGICAL APPROACHES

Surgery-Related Anatomy

The anterior cranial fossa is bound anteriorly by the frontal bone, inferiorly by the orbital roofs, the greater wing of the sphenoid posterior laterally, and the lesser sphenoid wings posterior medially (Fig. 5). The foramen cecum is in the midline behind the frontal bone, its posterior and lateral boundaries formed by the ethmoid bone. The crista galli is an osseous ridge arising from the midline of the ethmoid bone. The planum sphenoidale and the lesser wing of the sphenoid constitute the posterior floor of the anterior cranial fossa behind the cribiform plate.

The dura mater over the cribiform plates is thin. The blood supply to the anterior cranial fossa is from the anterior and posterior ethmoidal, middle meningeal, and ICA via the ophthalmic artery. The medial portion of the anterior cranial fossa contains the pneumatized frontal, ethmoidal, and sphenoid sinuses.

Fig. 5. Drawings providing an overview of the anterior cranial fossa anatomy and the surgical approaches.
The roof of the orbit is formed by the orbital process of the frontal bone, whereas the greater and lesser wings of the sphenoid form the apex and lateral wall of the orbits. The zygomatic bone marks its lateral wall. The lacrimal, ethmoid, palantine, and sphenoid bones comprise the medial wall of the orbit. The pterygoid plate lies beneath the greater wing of the sphenoid. The foramen rotundum and the inferior orbital fissure lie at the anterior end of the pterygoid plates. At the posterior end lies the foramen ovale.

The ethmoidal foramina are situated at the level of the frontoethmoidal suture. These foramina transmit the ethmoidal arteries and nerves. The posterior ethmoidal foramen is approximately 5 mm anterior to the orbital opening of the optic canal. The optic foramen lies at the posterior limit of the anterior cranial base.

**Indications for Surgery and Surgery-Related Principles**

Our indications for surgery and its principles are based on the experience of the senior author (T.F.) with 76 cases of malignant tumors of the anterior skull base. Lesions in this series were giant pituitary carcinomas (two cases), malignant craniohypophysealoma (one), malignant meningioma (six), teratocarcinoma (one), esthesioneuroblastoma (13), plasmacytoma (two), hemangiopericytoma (four), ethmoid small cell carcinoma (four), rhabdomyosarcoma (two), liposarcoma (one), basal cell carcinoma (one), malignant ameloblastoma (two), lacrimal gland carcinoma (one), squamous cell carcinoma (10), adenoid cystic carcinoma (four), chondrosarcoma (four), chordoma of the upper clivus (12), giant cell tumor (two), and adenocarcinoma (four cases).

In our experience, surgery should be reserved for cases in which at least 2 to 5 years of useful life remain following aggressive resection. Bilateral cavernous sinus involvement is a contraindication to surgery. To ensure patient longevity cranial nerve sacrifice, orbitectomy, eye exenteration, unilateral cavernous sinus sacrifice, or CA bypass may be necessary. These are only justified if they lead to complete en bloc tumor resection.

Surgical approaches to the anterior cranial fossa can be divided into five categories: 1) unilateral frontal transbasal; 2) bilateral transbasal; 3) extended bifrontal transbasal; 4) combined transbasal transcavernous; and 5) transfacial–transbasal. The basic principle underlying each approach is en bloc resection of the tumor. A goal of a tumor-free margin of 1 cm is attempted. Because of the propensity for spread of malignant cells, quick debulking is needed. The use of suction systems such as the Fukushima suction retractor help prevent spread of malignant cells. The drill is used minimally with a surgical osteome. After the scalp is reflected anteriorly, the long vascularized pericranial flap will be raised, with care taken not to lacerate or perforate the pericranium.

**Unilateral and Bilateral Transbasal Approach.** The unilateral and bilateral transbasal approaches begin with a bicoronal skin incision, uni- or bifrontal craniotomy, opening of the frontal sinuses, and a subfrontal extradural approach to lesions. Whereas a unilateral approach may suffice for lateral anterior fossa lesions, middle anterior fossa lesions require a bifrontal approach.

The patient is positioned supine in a Mayfield head holder with the neck slightly flexed and the vertex up. One of three variations of the bicoronal incision can be used (Fig. 7). Over the temporalis muscle, the areolar tissue and the superficial fat pad are raised with the flap to preserve the frontalis and zygomatic branches of the facial nerve. Over the vertex toward the forehead, the scalp is elevated strictly as the subgaleal flap, leaving the loose connective tissue attached to the pericranium. The pericranium is preserved between both temporalis muscles.

One centimeter from the orbital edge, the pericranial flap is raised with the frontalis skin flap to expose the orbital ridge. Care must be taken to avoid injury to the supraorbital nerve. The supraorbital nerve may be freed from the supraorbital foramen by using a drill or osteome. After the scalp is reflected anteriorly, the long vascularized pericranial flap will be raised, with care taken not to lacerate or perforate the pericranium. This graft can extend several inches behind the scalp incision.

The anteromedial portion of the temporal muscle is elevated posteriorly approximately 1 to 2 inches to expose the orbitotemporal rim. The first step in the craniotomy is a bifrontal craniotomy followed by the removal of the supraorbital rim (Fig. 8). Six burr holes are made using an 11-mm perforator or a 5-mm round burr. The initial two
burr holes are made through the orbitotemporal region just above the pterion, and the other two burr holes are placed just above the supraorbital foramen. The last two burr holes are made in the midline: one burr hole over the sagittal sinus 1 inch anterior to the Bregma and the other just above the frontal sinus. A bifrontal osteoplastic craniotomy is then performed.

**Extended Bifrontal Transbasal Approach.** The extended bifrontal transbasal approach adds an orbitofronto-ethmoidal osteotomy to the transbasal approach described by Derome.\(^2^{1-23,36}\) This approach can be used to excise midline tumors involving the clivus and sphenoid sinus and extending from the base of the dorsum sellae to the foramen magnum, with some involvement of the petrous apices. Complete loss of olfactory sensation and resultant loss of taste sensation can usually be expected after surgery via this approach. A technique for olfactory nerve preservation, as well as that of the cribriform plate, has been described.\(^72\)

After bifrontal dural elevation, a sagittal saw is used to remove the supraorbital bar. A midline osteotomy is made at the nasion, just superior to the medial canthal ligament. The peri-orbita is freed from the orbital roof. While protecting the periorbita with a brain spatula, the midline frontal bone and the orbital roofs are cut approximately 1 cm deep to the orbital ridge by using a saw or osteotomes. The lateral orbitotemporal groove is cut using a coarse diamond drill to detach the lateral orbital rim. Finally, the lateral orbital wall and the ridge are cut from the zygoma.

Following the removal of the supraorbital bar, the frontal basal dura is further separated from the anterior skull base. Using a periosteal elevator or a Fukushima dural elevator, the dura is elevated bilaterally, medial to the sphenoid ridge. The midline basal dura is dissected from the olfactory groove and from the crista galli. This olfactory portion of the dura should be sutured closed in a watertight fashion to prevent a CSF leak. It may be necessary to use a small tissue graft in patients in whom the dura is thin over the ethmoids.

The orbital roof is removed using a rongeur or Kerrison punch. The optic canal may be unroofed if the surgery is expected to extend to the posterior edge of the anterior fossa or if there is involvement of the anterior clinoid process. The diamond drill must be constantly cooled to avoid overheating the optic nerves. The medial orbital wall may also be resected. The key element of the frontal transbasal approach is the removal of the ethmoid and sphenoid sinuses. Once this is done, the clivus, sella floor, carotid groove, posterior pharyngeal mucosa, nasal septum, and the superior turbinates are visualized (Fig. 9).

**Combined Transbasal Transcavernous Approach.** In cases of cavernous sinus involvement, the transbasal approach can be combined with a transcavernous approach. This involves addition of an orbitozygomatic osteotomy. The anterior clinoid process is removed extradurally.\(^53\) The tuberculum sellae and the planum sphenoidal can be removed using a diamond burr to open widely the sphenoid sinus and the ethmoid sinus. Drilling may continue anteriorly in the midline to open the ethmoid sinus after removing the posterior cribriform plate. After removal of the anterior clinoid process, the anterior bend of the C\(_1\) segment of the ICA can be observed. The contralateral C\(_1\) can be observed within the sphenoid sinus. Using diamond burrs, the lateral wall of the superior orbital fissure can be removed. We perform the extradural temporopolar approach to free the cavernous sinus. The dura propria of the anterior medial temporal pole is elevated from the true cavernous membrane. This dissection begins by cutting the dura in the superior few millimeters of the superior orbital fissure. The dissection is carried down to the V\(_2\) segment. The dura propria is raised from V\(_2\) segment and the adjacent gasserian ganglia. With continuous retractor pressure on the temporal lobe, the dura continues to separate toward the petrous ridge. Dural elevation medially is limited by the tentorial edge. The posterior limit will be the V\(_3\) branch of the trigeminal nerve as it exits the ganglion. The oculomotor, trochlear, and trigeminal nerve branches (V\(_1\) and V\(_2\)) can be seen through the thin veil of the true cavernous membrane. The dura is opened and the anterior proximal 1 to 2 cm of the sylvian fissure arachnoid is split using sharp dissection.

Once the anteromedial and -lateral cavernous sinus area is exposed, attention is focused on exposing the posterior cavernous sinus region. The dura is elevated from the subtemporal middle fossa to the point where it is tethered by its attachment to the foramen ovale. The middle meningeal artery and its accompanying vein are coagulated and divided to allow full dural elevation from the middle fossa. Venous hemorrhage deriving from the middle fossa over the trigeminal nerve is controlled using Surgicel. Coagulation of these veins, which communicate with the
pterygoid venous plexus, will result in facial dysesthesias. The entire subtemporal middle fossa floor is shaved flat using a coarse diamond to obtain optimal visualization. The dura is elevated from the middle fossa to identify the arcuate eminence, the geniculate ganglion, and the greater superficial petrosal nerve. The foramen ovale is totally unroofed to expose the peripheral branches of V3 as it passes into the infratemporal area. This dissection is necessary to mobilize V3 and to elevate the gasserian ganglion to achieve anterior translocation of the trigeminal complex. The intrapetrous ICA is exposed by removing the bone over the posterolateral (Glasscock) triangle. With anterior translocation of the posterior border of V3, the “rhomboid area” can be identified. This is delineated by the greater superficial petrosal nerve, geniculate ganglion, arcuate eminence, the petrous ridge, and the posterior edge of V3. The petrous apex beneath the rhomboid can be safely removed medial to the C6 petrous CA and anterior to the arcuate eminence. This will expose the internal auditory canal, the posterior fossa dura, and the clivus.

The dural opening is continued laterally. The dura remains tethered at the fibrous dural rings that encase the CA and the cranial nerves. The external ring around the CA is cut, freeing that structure. The falciform ligament is cut, freeing the optic nerve. The tentorial edge is composed of attachments to the petrous tip, as well as the anterior and the posterior clinoid processes. The fibrous ring over the oculomotor nerve as it passes through the porus oculomotorius is also opened. The dura over the trochlear nerve is opened. If necessary the posterior clinoid process and the dorsum sellae can be removed using a diamond drill. Dissection is accomplished by drilling to the posterior clinoid process and the dorsum sellae. The cavernous sinus can be entered via one of the entry corridors initially depicted by Parkinson51 and subsequently summarized in the scheme proposed by Fukushima and Day.28

**Reconstruction of the Anterior Skull Base and Closure.** The supraorbital bar is replaced and fixed in place by using a plating system. The vascularized pericranial graft is laid over the supraorbital bar to the clivus. The edge of the pericranial flap is sutured to the posterior pharyngeal soft tissue, tuberculum sellae, and orbital fascia. The frontal basal dead space is filled with a free fat graft harvested from the abdomen. A watertight dural closure is needed. A microplate is used to replace the frontal bone flap. A lumbar drain is placed for 24 hours postoperatively to minimize the risk of postoperative CSF leakage.

**Transbasal–Transfacial Approach.** The transbasal approach can be combined with the transfacial approach for complete resection of anterior fossa malignancies that are contiguous with paranasal sinuses. Variations of the transfacial approach include the lateral rhinotomy and midfacial degloving.64

The lateral rhinotomy incision is made on the side of greater tumor involvement. The incision begins medial to the medial eyebrow, extended along the nasofacial groove and around the alar rim into the nasal vestibule. If palatal resection is necessary, the incision can be extended to split the lip in the midline.

A lateral osteotomy is made through the frontal process of the maxilla. Care is taken not to injure the nasolacrimal duct or the ligament of the medial canthus. Extension of the osteotomy across the roof of the nose allows reflection
of the nasal skeleton and overlying soft tissue to the contralateral side. The osseous nasal septum and posterior septal cartilage are separated posteriorly from the nasal bones and external nasal structure. The lacrimal duct is transected. The periorbita is elevated from the medial and inferior surfaces of the orbit, the lamina papyracea, and maxilla.

Osteotomies are created through the ipsilateral inferior meatus, medial orbital floor, and the frontoethmoid suture anterior to the optic foramen. Following the osteotomies, the tumor can be resected en bloc via the transcranial and transfacial openings.

In the midfacial degloving technique the midfacial soft tissues are reflected superiorly. Bilateral exposure of the ethmoid and maxillary area is obtained. To free the midface soft tissue, bilateral sublabial incisions, intranasal transfixation incision, bilateral intercartilaginous incisions between the upper and lower cartilages, and bilateral piritiform aperture incisions extending to the vestibule of the nose are made. Following elevation of the soft tissue, Penrose drains are inserted into both nares and used to retract the midfacial soft tissue superiorly to the level of the glabella and orbits. Osteotomies are performed as previously described via the lateral rhinotomy approach.

The pericranial graft is used to provide a vascularized surface. The nasal tip is returned to its normal position. The columellar, intercartilaginous, and sublabial incisions are closed using absorbable sutures. A nasal splint is placed over the nasal dorsum.

Complication Avoidance

Both CSF leakage and meningitis are potential complications that can be minimized by obtaining a watertight dural closure. A vascularized pericranial graft to exclude the cranialized frontal sinus is critical. Significant wound complications are rare. Cranial nerve deficits may occur, but with meticulous microsurgical technique these too can be avoided.

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