Complex Skull Base Reconstructions in Kadish D Esthesioneuroblastoma: Case Report

Sheri K. Palejwala¹ Saurabh Sharma² Christopher H. Le² Eugene Chang² Audrey B. Erman² G. Michael Lemole, Jr¹

¹Division of Neurosurgery, Department of Surgery, University of Arizona, Tucson, Arizona, United States
²Department of Otolaryngology, University of Arizona, Tucson, Arizona, United States


Abstract

Introduction Advanced Kadish stage esthesioneuroblastoma requires more extensive resections and aggressive adjuvant therapy to obtain adequate disease-free control, which can lead to higher complication rates. We describe the case of a patient with Kadish D esthesioneuroblastoma who underwent multiple surgeries for infectious, neurologic, and wound complications, highlighting potential preventative and salvage techniques.

Case Presentation A 61-year-old man who presented with a large left-sided esthesioneuroblastoma, extending into the orbit, frontal lobe, and parapharyngeal nodes. He underwent margin-free endoscopic-assisted craniofacial resection with adjuvant craniofacial and cervical radiotherapy and concomitant chemotherapy. He then returned with breakdown of his skull base reconstruction and subsequent frontal infections and ultimately received 10 surgical procedures with surgeries for infection-related issues including craniectomy and abscess evacuation. He also had surgeries for skull base reconstruction and CSF leak, repaired with vascularized and free autologous grafts and flaps, synthetic tissues, and CSF diversion.

Discussion Extensive, high Kadish stage tumors necessitate radical surgical resection, radiation, and chemotherapy, which can lead to complications. Ultimately, there are several options available to surgeons, and although precautions should be taken whenever possible, risk of wound breakdown, leak, or infection should not preclude radical surgical resection and aggressive adjuvant therapies in the treatment of esthesioneuroblastoma.

Keywords
► esthesioneuroblastoma
► olfactory neuroblastoma
► skull base reconstruction
► cerebrospinal fluid leak
► pneumocephalus
► sinonasal malignancy
► complications

Introduction Esthesioneuroblastoma, or olfactory neuroblastoma, was first described by Berger et al, in 1924 as l’esthésioneuroépithéliome olfactif.¹ Since then several hundred patients have been described in case series, single-institution reviews, and meta-analyses.²–26 Patient presentation is often nonspecific, including congestion and sinusitis-like symptoms, which makes diagnosis challenging, such that most patients have advanced disease at the time of diagnosis.¹5,18,27 The gold standard for the treatment of esthesioneuroblastomas is craniofacial resection with histologically proven disease-free margins, with the use of radiotherapy.²,7,8,10,12,20,25 The use of chemotherapy, neoadjuvant therapy, neck dissection, and irradiation, however, remains controversial.⁵–⁷,11,14,22

Esthesioneuroblastoma has an established propensity for being locally aggressive with the possibility of distal metastases and leading to decades-delayed recurrences.⁵,¹¹,12,20 In
an effort to prolong disease-free survival, most advocate for radical resection with clear margins, which has been shown to double disease-free survival, especially in the setting of advanced Kadish tumors.2,5,7,28 Additionally, greater tumor extent often necessitates substantial resections, which have both been associated with complication rates as high as 33%.28–30 More extensive disease, meticulous surgical resection, and aggressive adjuvant therapies together increase the likelihood of adverse events, including cerebrospinal fluid (CSF) leak, neurologic deficits, and infectious complications.

We performed a retrospective chart review to describe the history of a single patient with Kadish D esthesioneuroblastoma, who underwent endoscopic-assisted craniofacial resection followed by adjuvant radiation and chemotherapy. His course was complicated by multiple infections and CSF leaks necessitating several skull base reconstructions. We describe his case to elucidate the multiple and successive escalations for reconstruction available to skull base surgeons, even in the face of malignancy, radiation, chemotherapy, and infection.

**Case Report**

A 61-year-old man, with a several decade history of chronic sinusitis, presented with a 5-month history of nasal congestion, decreased sense of taste and smell, and intermittent yellow nasal drainage, which failed to resolve with antibiotics, scant epistaxis, left ptosis, and medial periorbital edema. Nasal endoscopy revealed a 7-cm nasal mass, and biopsy demonstrated esthesioneuroblastoma. On imaging there was a homogenously enhancing, erosive left skull base lesion with extension into the right nasal cavity, bilateral ethmoid, left maxillary, frontal, and sphenoid sinuses with anterior cranial fossa extension and leptomeningeal enhancement (Fig. 1). Two mildly enlarged fludeoxyglucose (FDG)–avid left parapharyngeal lymph nodes were seen on positron emission tomography–computed tomography (PET-CT).

He subsequently underwent a combined level 1 transbasal and endoscopic endonasal approach for complete, margin-free resection. At the start of the procedure, the bilateral middle and superior turbinate were removed, and tumor was resected from the nasal cavity, ethmoid and sphenoid sinuses, and frontal recess. This was followed by a transbasal level 1 craniotomy with removal of the orbital bar and both intra- and extradural tumor resection and resection of the skull base dura. All margins were negative and reconstruction was performed with a large vascularized pericranial flap onlay, four standard and two firm Nopores (Polyganics, the Netherlands), and nasal trumpets. The patient was discharged home on postoperative day (POD) 5 without any perioperative complications.

The patient then underwent intensity-modulated radiation therapy (IMRT) with 54 Gy to the tumor bed and 70 Gy for his cervical lymph node disease, with two cycles of cisplatin and etoposide concurrently. After completing therapy, 3.5 months after surgery, he had no radiographic evidence of residual or recurrent disease (Fig. 2). PET-CT 6 months after surgery revealed decreased size and activity of the parapharyngeal nodes as well. His postoperative course was complicated by radiation-induced dysphagia, treated with temporary percutaneous endoscopic gastrostomy (PEG) tube placement, and nasal crusting, managed with periodic debrides and antibiotics when indicated.

He was regularly followed by the neurosurgical, otolaryngology, radiation, and medical oncology teams. Nearly 1 year after surgery, he presented with progressive left eye ptosis and painful forehead and left canthal lesions. Magnetic resonance imaging (MRI) showed avid enhancement and a neofrontal sinus in the epidural space (Fig. 3). He was taken to the operating room 15 months after his initial resection for endoscopic debriement, biopsies to rule out intranasal recurrence, and fine-needle aspiration (FNA) of the forehead masses. He was found to have significant mucopurulence, osteoradione-rosis of portions of his frontal bone, which were then removed, and shrinking of his pericranial flap with an airspace between the pericranium and frontal bone (Fig. 4). He then underwent FNA of the forehead mass and left endoscopic dacryocystorhinostomy (DCR). No malignancy was found, but forehead cultures grew *Proteus mirabilis* and *Klebsiella pneumoniae* for which he was treated with intravenous (IV) metronidazole and ceftriaxone for 8 weeks. After extensive multidisciplinary discussions, the decision was made to proceed with removal of the necrotic bone flap.

Seventeen months after initial resection, the patient underwent craniectomy, irrigation and debridement, titanium mesh cranioplasty, and skull base reconstruction with an autologous fascia lata graft. There was no gross
extracranial infection, but an obvious epidural collection with fibrinous exudate, and osteomyelitis and necrosis of the orbital bar. Both were removed and discarded. We also encountered a focal dehiscence in the pericranium, allowing communication between the endonasal and intracranial spaces. This was repaired with a water-tight closure using a fascia lata autograft. After extensive irrigation and debridement, cranial reconstruction was performed with titanium mesh, which was contoured to the nasal bridge and provided medial orbital bar reconstruction. No evidence of malignancy was noted either grossly or on histopathology.

He was discharged home on POD 5 with 4 weeks of IV metronidazole and ceftriaxone. Cultures revealed Mycobacterium chelonae for which he was treated with additional IV amikacin, IV tigecycline, and PO clarithromycin for an additional 2 weeks, and then continued on PO doxycycline and clarithromycin for 12 weeks.

Four months later, CT revealed a persistent intracranial air pocket with an area of presumed continued nasal-intracranial communication. After completing his antibiotic regimen, the patient underwent cranial reconstruction with a custom polyetheretherketone (PEEK) bone flap with an orbital bar extension and vascularized left radial forearm free flap, to cover the anterior skull base defect, 5 months after the craniectomy (22 months after initial surgery). Reconstruction required partial takedown of the pericranial flap laterally to access the skull base posteriorly to the level of the planum sphenoidale. We encountered a small area of CSF leak around the left orbit that was reconstructed using primary closure and Tisseel fibrin sealant (Baxter International Inc., Deerfield, Illinois, United States). The vascularized radial forearm myofascial flap was anastomosed to the superficial temporal artery and vein and was found to be well perfused and laying freely. It was secured to the dura and skull base, and nasal endoscopy confirmed complete skull base closure. Postoperative imaging is seen in Fig. 5.

On POD 2 the patient experienced acute altered mental status and disinhibition, CT revealed a large bifrontal epidural hematoma, and the patient was taken emergently to the operating room for evacuation (Fig. 6). Another small CSF leak was noted and also repaired with primary closure, Tisseel fibrin sealant, and placement of a lumbar drain. His mental status gradually returned to baseline and he was discharged home on POD 7.

He returned to the emergency department 2 days after discharge after reports of strange disinhibited behavior. CT revealed intracranial air with a new endonasal-intracranial focus of communication, suspicious for tension pneumcephalus (Fig. 7). He was taken to the operating room for yet another skull base reconstruction where the new defect was clearly noted. An AlloDerm (LifeCell Corp., Woodlands, Texas,
United States) graft was cut to size and sutured both to the planum sphenoidale and the peri-orbita bilaterally. Additional AlloDerm was used as an on-lay over the friable frontal lobe dura and secured with Evicel fibrin sealant (Johnson & Johnson Wound Management, Somerville, New Jersey, United States). The myofascial flap was reapproximated and reconstruction was again confirmed with nasal endoscopy (►Fig. 8). He had a gradual return to his intact neurologic baseline and was discharged home on POD 9.

The patient returned 1 month later to the emergency department (ED) with complaints of 2 days of severe progressive headache and subgaleal fluid collection. He was treated with 5 days of CSF drainage via a lumbar drain and discharged on hospital day 8, after observation for two days after drain removal. Unfortunately, he returned with recurrence of the subgaleal fluid collection, and moderate amount of fluid expressed from a forehead pustule. Imaging revealed increase in the subgaleal and epidural fluid, and he was once...
again admitted for an epidural-peritoneal shunt, without immediate perioperative complications. However, the epidural fluid culture later returned as *Enterobacter cloacae* and was treated with 8 total weeks of IV meropenem. He also complained of abdominal pain concerning for infectious or chemical peritonitis, while abdominal CT revealed fat infiltration and a right lower quadrant fluid loculation. He was reluctant to have his shunt removed due to concern for fluid reaccumulation and possible subsequent infection.

Seven weeks after shunt placement, the patient presented to the ED with rhinorrhea and pneumocephalus and was taken for shunt removal and epidural drain placement. Four days later, he underwent yet another skull base reconstruction with repositioning of the still viable myofascial flap, additional buttressing with an abdominal fat graft, and insertion of lumbar drain. There was no evidence of infection or tumor recurrence. His postoperative course was complicated with abdominal hematoma that was evacuated at bedside, and he was discharged home on POD 8.

After the aforementioned 8 weeks of IV meropenem, the patient was transitioned to 4 weeks of PO levofloxacin and continued to be monitored with imaging and clinical nasal endoscopy. MRI performed 26 months after initial resection then revealed right temporoparietal and falcine nodular dural enhancement concerning for metastatic disease (Fig. 9). PET-CT revealed progression of his intracranial disease and pulmonary nodules. He was referred to both medical and radiation oncology for salvage adjuvant treatment.

**Discussion**

The first, and most commonly used, staging system for esthesioneuroblastoma was described in 1976 by Kadish et al and was later modified by Foote et al to include involvement of cervical lymph node and distant metastases. High Kadish classification, especially with cervical lymph node involvement, is known to be a profound negative prognosticator, cutting survival in half and increasing the rate of distal...
metastasis by 35%. On the other hand, aggressive surgical resection, with clear margins, is a strong predictor of disease freedom. Anterior skull base tumors with positive margins were found to have double the incidence of local recurrence and half the survival of complete resections. The widespread use of craniofacial approaches has improved our ability to obtain complete resections with histologically disease-free margins, even in the setting of extensive, high Kadish stage tumors. Craniofacial resection has been shown to increase progression-free survival from 37.5 to 82%.

Despite the favorable outcomes with extensive resection, single-modality treatment, including surgery in isolation, led to poor results and higher rates of both local recurrence and distal metastatic disease. Besides radical resection, the highest control rates were found with the addition of radiation therapy. Dulguerov et al performed a meta-analysis of 390 patients which ultimately showed the best outcomes in esthesioneuroblastoma occurred when margin-free resection was followed by radiotherapy; this has also been corroborated by several large studies. Although controversial, many groups argue for the addition of chemotherapy as well, to achieve the longest duration of tumor freedom. Multi-modality treatment is especially recommended in those with high Kadish stage esthesioneuroblastoma.

Although aggressive management of these persistent malignant tumors is advocated, there are treatment effects and complications that should be considered. Postoperative complication rates for craniofacial approaches can affect one in three patients undergoing anterior craniofacial resection based on an International Collaborative Study, with wound complications in 18% of patients. Although there have been significant advances in skull base reconstruction, especially from endoscopic approaches, CSF leak still remains a significant concern. Some studies advocate the use of synthetics, whereas others use vascularized nasoseptal flaps, and still other groups encourage the use of gasket seals. Ultimately there are numerous options available to the surgeon, which can be used simultaneously or even sequentially, should the need arise. Unfortunately, prior radiation treatment and wide intracranial tumor extension increase the risks of complications, including infection and leak.

Patients requiring postoperative radiation therapy will likely also incur decreased vascularization and increased fibrosis. Over time, most free flaps, without vascularization, will reabsorb, and these flaps can be salvaged by buttressing them with vascularized flaps from the outset. Additionally, in the setting of radiation, allografts have been shown to have a high rate of extrusion. Pedicled pericranial flaps are the most frequently used and easily accessible flaps and have been proven successful. Nasoseptal flaps have also been shown to have tremendous success, even in the setting of wide extensive tumors. However, in the setting of malignancy, they can only be safely used if histologically confirmed to be tumor-free. Unfortunately, in our case, this was not enough to prevent breakdown, with subsequent nasal-intracranial communication and an epidural, neofrontal sinus. Pedicled myocutaneous and myofascial flaps have also been used with success, including the pectoralis major, latissimus dorsi, and trapezius muscles. They have both the ability to fill dead space, dampen the effect of CSF and brain pulsations, and have a rich vascular supply. Although this provided a good salvage reconstruction for our patient, it was not without its own complication, a symptomatic epidural hematoma requiring emergent evacuation. Free tissue transfer and synthetic allografts are also available options, particularly when supplementing more robust vascularized flaps, in the setting of chemotherapy and radiation. This was also corroborated with our own series and experience with this patient.

**Conclusion**

Studies have shown that greater tumor extent, especially in the setting of intracranial extension, cervical lymph node involvement, and distal metastases are associated with greater rates of recurrence and poorer prognoses.
In light of this, initial treatment, both medical and surgical, should be aimed at curing the primary disease. Measures should be taken to ensure adequate skull base reconstruction from the outset, as well as any risk mitigating actions that can be performed. Nevertheless, sequential escalation of varied repair techniques is possible and often necessary in these challenging cases. Early use of vascularized flaps should be considered after aggressive skull base resections, radiation, and chemotherapy. Ultimately, there are several options available to surgeons, and although precautions should be taken whenever possible, risk of wound breakdown, leak, or infection should not preclude radical surgical resection and aggressive adjuvant therapies in the setting of malignancy.

Disclosures

The authors have no disclosures or conflicts of interest.

References