Management of Acute Facial Nerve and Parotid Injuries

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Abstract

Acute soft tissue trauma to the head and neck is a common reason for emergency department presentation and should be appropriately evaluated by a facial plastic surgeon. The evaluation of a patient who has suffered facial trauma should always include a comprehensive facial nerve exam and carry a low threshold of suspicion for parotid duct injury when involving the cheek. Injuries to the facial nerve and parotid duct can result in significant long-term functional, cosmetic, and emotional morbidity, particularly when diagnosis is delayed. In the repair of facial nerve transection, neurorrhaphy technique is primarily based on the ability to obtain tension-free anastomosis and outcomes are in large part related to timing of repair. Parotid duct injuries are generally repaired based on the site of ductal injury. In this article, we present a guide to the relevant anatomy of the facial nerve branches and the parotid duct, the important factors guiding treatment decisions alongside their related risks and benefits, as well as the management of complications of facial nerve neurorrhaphy and parotid duct injuries and repair.

Keywords
► facial trauma
► facial nerve
► parotid duct
► parotid gland

Acute soft tissue injuries to the head and neck are a common reason for patients to present to the emergency department and require evaluation by a facial plastic surgeon. Facial anatomy is complex, as it houses four of the five major senses as well as speech. Our face is the primary means by which others identify us as individuals as well as how we express ourselves to the world. The cheek is the subunit of the face with the largest surface area and is frequently injured. It is also the subunit overlaying the main trunk and inferior branches of the facial nerve (FN), the parotid gland, and the parotid duct (PD). While cutaneous soft tissue injuries to the cheek are easily identified on inspection, a high degree of suspicion must be maintained during the initial evaluation to determine if there has been damage to these deeper structures. Failure to identify or completely repair injuries to the FN or the parotid can lead to significant morbidity for the patient down the road.

Traumatic Facial Nerve Paralysis Secondary to Transection

The incidence of peripheral FN palsies ranges from 20 to 30 cases per 100,000 patients per year—one of the most common cranial nerve pathologies. While most cases are idiopathic, traumatic etiologies are responsible for up to 30% of cases, most in the form of temporal bone fractures. FN injury following soft tissue trauma usually occurs in the setting of stab wounds, gunshot wounds, or motor vehicle accidents.

Anatomy

The FN begins intracranially at the pons and then takes a complex course through the internal auditory canal into the temporal bone, where it makes two turns (“genus”) before it exits the skull base at the stylomastoid foramen. Peripherally,
the main trunk of the extratemporal FN courses anteriorly and deep to the posterior belly of the digastric for approximately 1 to 1.5 cm before it splits at the pes anserinus (Latin for “goose foot”) where it bifurcates into the temporofacial and cervico-facial divisions. Though the branching pattern of the distal FN is variable, the temporofacial branch typically divides into the terminal temporal (frontal) and zygomatic branches, while the lower cervicofacial branch trifurcates into the buccal, marginal mandibular, and cervical branches.

Injuries at different anatomical levels of the FN can produce different deficits. The more distal an injury, the more motor and less sensory and parasympathetic deficits. Nonmotor deficits to be evaluated include xerophthalmia, nasal dryness, dysgeusia or ageusia, and xerostomia.

**Evaluation**

The evaluation of a facial trauma victim should always include a complete cranial nerve exam. Assessment of the FN should be done in a systematic fashion, as particular deficits can elucidate the branches compromised. This should always be done prior to the injection of local anesthetic. The face is typically divided into five motor segments assessed in a superior to inferior direction: forehead, periorcular, midface, peripheral, and cervical. Assessment of the forehead involves brow elevation and brow furrowing. Periorcular evaluation focuses heavily on gentle and forceful eye closure to assess all muscular components of the orbicularis oculi. Lower eyelid position and tension is noted, but can be affected by midfacial function and ptosis as well. Midfacial function is assessed with wrinkling of the nose and smiling or lip elevation. The perioral function is evaluated by puckering and pouting motions and the cervical division by flexing of the platysma. Areas of obvious trauma can quickly be assessed with their relation to the lateral canthus. If the injury is medial to the lateral canthus, immediate repair may not be warranted as the nerve is more likely to regain function without surgical management due to extensive cross innervation. A distinction should be made between paresis and paralysis, as paretic muscles imply continuity of the nerve and such injuries can be managed conservatively.

**Timing of Exploration**

The presence of complete paralysis at the time of evaluation portends a poor prognosis as compared with delayed paralysis. In such scenarios, the timing of exploration and repair has consistently been the largest predictive factor to successful outcomes in traumatic FN transection. Exploration should occur within 72 hours of injury to allow electrical stimulation of distal branches to facilitate identification, otherwise Wallerian degeneration would progress to where stimulation is not feasible. Electroneuronography can be useful between 3 and 21 days after injury but starting at 3 weeks electromyography (EMG) is the gold standard to determine the potential for reinnervation. EMG begins to demonstrate fibrillation potentials 2 to 3 weeks after denervation of the facial musculature, and subsequent polyphasic action potentials (PAPs) indicate regeneration is occurring. If PAPs are present, this may indicate a higher likelihood of spontaneous return of motor function, allowing for more conservative management. Absence of fibrillation potentials and PAP’s indicates irreversible fibrosis. During the first 3 months after injury, contraction secondary to scarring reduces the cross-sectional area of the nerve up to 70%.

In the cases of either delayed diagnosis or delayed intervention due to the patients’ instability, there is no definitive agreement regarding the optimal time for reinnervation procedures, though it is widely believed that the earliest repair possible results in the greatest functional outcome. Immediate direct neurorrhaphy has shown the best functional results. The current literature is contradictory, with some studies supporting better functional outcomes when repaired within 90 to 180 days, while others comparing immediate and early repair, defined as 6 to 18 months post injury, have demonstrated comparable outcomes. The latter study did show that nerve function returns significantly faster in the early repair arms. There is a large consensus that the best possible functional outcome after complete transection of the FN, regardless of the method of coaptation and time until intervention, is House-Brackmann grade III.

**Nerve Repair Techniques**

**Direct Neurorrhaphy**

Direct neurorrhaphy is the gold standard for FN repair. However, this technique requires a tension free neurorrhaphy of the severed nerve ends, which is not always possible. If tension is present at this site, impaired vascular flow may lead to scarring and inhibit axonal regeneration. Attempts can be made to mobilize the nerve inferiorly and superiorly to ease tension.

The neurorrhaphy itself is typically performed using an end-to-end coaptation following conservative trimming of the nerve ends. A cuff of epineurium is retracted away from the cut end to allow the fascicles to be trimmed sharply. The epineurium is then released to overlap the trimmed fascicles slightly, allowing the fascicles to lay gently end-to-end once the epineurium is approximated. The coaptation is done with 9-0 or 10-0 nylon sutures through the epineurium under an operating microscope (Fig. 1). Direct neurorrhaphy results in the best outcomes compared with other techniques in terms of both a faster and more significant improvement in House-Brackmann scores.

![Fig. 1](image-url) Intraoperative view following primary neurorrhaphy of transected facial nerve due to a penetrating cheek injury.
Difficulty identifying the proximal and distal ends of the FN in the setting of robust soft tissue injury can be problematic. Identifying the main trunk of the FN either within the mastoid or utilizing the reliable landmarks at its exit from the stylomastoid foramen may be helpful. Distal identification may require identification of peripheral branches with retrograde dissection toward the main trunk.

Peripherally, certain anatomical landmarks are helpful to approximate the location of each branch of the FN.

**Frontotemporal Branch**
The path can be estimated by Pitanguy’s line, which is a line drawn through two points: (A) midpoint between the root of the helix and the lateral canthus and (B) the root of the ear lobe.

**Buccal Branch**
It is located by its proximity to the PD, typically running within 1 cm inferior to the duct and along the same trajectory.

**Marginal Mandibular Branch**
Typically runs 1 to 2 cm inferior to the inferior border of the angle of the mandible, however, it can be found up to 3 cm inferiorly.

**Cervical Branch**
It is found that approximately 1 cm posterior to the gonion. Incisions 2 cm posterior to the gonion will typically preserve the inferior parotid pole and lower branches of the FN. Another approximation is a line drawn from the mentum to the mastoid process, with a perpendicular line 1 cm inferior to the angle of the mandible estimating the location of the nerve.

**Interposition Nerve Grafting**
If a tension free direct neurorrhaphy is not possible, an autograft or allograft can be used to bridge the gap and provide a tension free neurorrhaphy. Common autograft donor sites include the great auricular nerve (GAN), sural nerve (SN), and the medial antebrachial cutaneous (MABC) nerves.

**Great Auricular Nerve**
The GAN is utilized most frequently given its proximity, roughly equivalent cross-sectional area, and low donor site morbidity. The GAN can typically provide 6 to 8 cm in length. It is found approximately 6.5 cm below the inferior wall of the external acoustic meatus and 0.5 cm above the external jugular vein. Another method of identification is with a line drawn from the angle of the mandible to the mastoid tip, with the GAN bisecting this line as it passes over the sternocleidomastoid. It runs deep to the platysma muscle in the same general course of the external jugular vein.

**Sural Nerve**
The SN is harvested when length is a concern or there is a need to graft multiple distal nerve branches. The SN approximates the cross-sectional area of the FN and can supply up to 40 cm of graft. It is found as it leaves the tibial nerve in the popliteal fossa and descends between the two heads of the gastrocnemius, piercing the deep fascia at the posterior aspect of the lower leg. It is often just posterior to the lateral malleolus and can be harvested with either one long incision, several transverse “stair-step” incisions, or endoscopically. For open harvest, the first incision is made 2 cm posterior and 2 cm proximal to the lateral malleolus, as the nerve has not undergone significant branching at this point. If the small saphenous vein is encountered, the nerve lies just medial to it. A tendon stripper may be useful to clear off the soft tissue.

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**Fig. 2** The path of the frontal branch of the facial nerve can be estimated by Pitanguy’s line, which is a line drawn through two points: (A) midpoint between the root of the helix and the lateral canthus and (B) the root of the ear lobe.

**Fig. 3** The great auricular nerve is identified by perpendicularly bisecting a line drawn from the angle of the mandible to the mastoid tip over the sternocleidomastoid.
endoscopic harvest of the SN, a horizontal incision is made approximately 1 cm behind the lateral malleolus. The saphenous vein is identified, and the SN identified immediately posterior to this. The endoscope and endoharvester are passed along the length of the nerve to skeletonize the required length. The nerve is then cut proximally and delivered from the inferior incision (►Fig. 5).10 The disadvantages to using the SN as an autograft are the need for another operative site, scarring, and sensory deficits of the lateral foot.

Medial Antebrachial Cutaneous Nerve
The MABC nerve offers an ideal size match and branching pattern as an autologous donor nerve. This nerve can be used when repairing defects extending from the main trunk or intra-temporal nerve to multiple distal branches. A cutaneous incision is placed along the medial bicipital groove (►Fig. 6). The MABC nerve is located near the cephalic vein in the superior forearm, and can be found as it emerges laterally from the inferior aspect of the biceps brachii tendon as it pierces the inter-epicondylar line. At this level, it is approximately 1 cm medial to the cephalic vein and can be seen decussating to innervate the forearm skin11 (►Figs. 7 and 8).

Polarity of Grafts
Polarity is a commonly debated topic when grafting nerves. Some advocate reversing an autograft’s polarity under the belief that it reduces aberrant re-arborization of nerve fibers. Multiple animal studies have demonstrated that reversing the polarity of an autograft did not have significant effects on the total number of axons regenerated, nerve conduction velocity, or functional outcomes.12 More consideration should be given to nerve diameter matchup than polarity in optimal nerve reconstruction.

Nerve Allografts
If the patient’s GAN, SN, or MABC nerves are either unavailable or unable to be used, an allograft can be used to bridge the gap and provide a tension free repair. Situations in which an allograft may be preferred over an autograft are patients who have received radiation at the donor and/or recipient site, tumor invasion, or significant comorbidities precluding prolonged operative times. A major benefit of allografts is the lack of donor site morbidity. Traditional allografts have fallen out of favor due to the requirement for long-term host immunosuppression, leading to the development of decellularized nerve allografts.13 These allografts have reduced
immunogenicity but retain the beneficial characteristics of scaffolding and extracellular proteins that may facilitate nerve regeneration.14 Allografts performed for peripheral nerve repair support meaningful recovery of up to 89% in sensory nerves and 86% in motor nerves.15 While the majority of studies center around neurorrhaphy in the hand, recent data from reinnervation of the lingual and inferior alveolar nerves have demonstrated upward of 94% functional sensory recovery at 1 year postoperatively.12 The limited data regarding allografts in FN repair reveals suboptimal outcomes compared with direct neurorrhaphy, but allografts are still a feasible choice when an autograft is not an option (►Figs. 9 and 10).16

Complications and Management

Complications related to traumatic FN paralysis can occur as direct sequelae from the paralysis or indirectly related to the neurorrhaphy. Immediate complications related to nerve paralysis are due to the loss of both motor function and static muscle tone of the mimetic muscles of the face. A complete discussion regarding management of such complications is outside the scope of this chapter, but several of the more serious acute issues warrant mention. Loss of the upper branches of the FN can result in paralytic lagophthalmos. Corneal protection can be compromised with inability to close the eye resulting in corneal abrasions and ulcerations. Patients who suffer a concomitant injury to the ophthalmic division of the trigeminal nerve (V1) are at even higher risk of neurotropic keratopathy due to an insensate cornea. Immediate treatment of paralytic lagophthalmos consists of lubricating eye drops and a moisture chamber to mimic the physiologic function of a closed eyelid, with a low threshold to perform upper eyelid weight implantation or lateral tarsorrhaphy if the patient develops ocular pain or corneal irritation.

Complications related to neurorrhaphy include synkinesis, donor site morbidity such as pain, scarring, wound complications, and sensory deficits, and failure of meaningful functional recovery.

Synkinesis or Nonflaccid Facial Paralysis

Synkinesis is the involuntary contraction of a muscle associated with the voluntary contraction of another muscle. Synkinesis is thought to be secondary to aberrant axonal regrowth where the axons do not attach back to their original fascicles.3,17 When synkinesis occurs, it can begin as soon as 3 to 4 months after injury and can be quite distressing. The most common types of facial synkinesis include oral-ocular (involuntary eye closure during voluntary mouth movements) and ocular-oral (involuntary mouth movement during voluntary eye closure). The goals of treatment include improving resting oral commissure position, oral competence, eye closure, smile symmetry, and mimetic movement spontaneity.17 Treatment strategies include physiotherapy by an FN trained physical or occupational therapist, botulinum toxin injections, and surgical correction or augmentation. Physiotherapy aims to maximize voluntary control of the FN and associated mimetic muscles through exercise with the ultimate goal being to retrain the nerve to its proper functional pathways.18 Physiotherapy has shown to be effective in both therapy of active synkinesis as well as preventing nerve dysfunction following injuries. Botulinum toxin injections are best utilized as an adjuvant to physiotherapy. The goal is to selectively inhibit muscle contractibility of the affected areas. These are most commonly the orbicularis oculi, depressor anguli oris, mentalis, and the platysma. Choi et al demonstrated that when employed as an adjuvant treatment with physiotherapy, patients exhibited a better quality of life and improved appearance.19 Surgical treatment involves static correction of muscle hypofunction such as with brow lifting, or more dynamic effects with myectomies and selective neurolysis.18

Management of Acute Parotid Gland and Parotid Duct Injuries

Epidemiology

Parotid injuries typically occur following penetrating trauma to the cheek and are associated with damage to the adjacent
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Facial structures, such as the facial and lingual nerves, the ear, and bony structures of the face. PD injuries account for approximately 0.21% of all trauma cases. While approximately half of these cases are recognized as acute parotid trauma, the rest present as complications of an undiagnosed injury, such as sialoceles or fistulas.

**Anatomy**

The parotid gland overlies the ramus of the mandible and is bounded superiorly by the zygoma, anteriorly by the masseter, and posteriorly by the external auditory canal and the superior aspect of the sternocleidomastoid. The gland is divided into superficial and deep portions by the FN anatomically, and the retromandibular vein radiologically. As the PD emerges anteriorly from the parotid gland, it travels superficial to the masseter in close relation to the transverse facial artery and buccal branch of the FN. At the anterior border of the masseter, the duct penetrates the buccinator muscle (the “masseteric bend”) and oral mucosa before terminating at the level of the second maxillary molar. The path of the PD can be envisioned as following an “S-shaped” path that crosses an imaginary line which joins the tragus-antitragus point to the middle of the upper lip (►Fig. 11). Injuries involving this area must be carefully assessed for ductal trauma.

**Evaluation**

Any penetrating injury or deep laceration of the cheek along a line drawn from the tragus to the midportion of the upper lip has the potential to damage the parotid gland and/or the PD. The classification of parotid gland and ductal injuries according to location was described by Van Sickels as injuries posterior to the masseter (region A), injuries to the masseter (region B), and injuries anterior to the masseter (region C) (►Fig. 12; Table 1). Region A injuries occur in the gland only, region B corresponds to the duct as it runs superficial to the masseter, and region C corresponds to the duct anterior to the masseter as it passes through the buccal space and attaches to the buccal mucosa opposite to the second maxillary molar. It is important to remember that the buccal branch of the FN parallels the PD, and concomitant injuries can occur in regions B and C (►Fig. 13). When an injury to the gland or duct is suspected, most clinicians advocate for repair within 24 hours after the injury prior to edema making ductal cannulation more tedious.

**Management**

Parotid Gland Injury

Isolated injury to the parotid gland (region A) is best treated with direct suturing of the enveloping fascia followed by application of a pressure dressing for 48 hours to reduce the
Table 1  Regions of parotid injury and associated complication rates

<table>
<thead>
<tr>
<th>Region of injury</th>
<th>Relation to masseter</th>
<th>Complication rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Posterior</td>
<td>22</td>
</tr>
<tr>
<td>B</td>
<td>Overlying</td>
<td>80</td>
</tr>
<tr>
<td>C</td>
<td>Anterior</td>
<td>80</td>
</tr>
</tbody>
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Source: Adapted from Gordin et al. 36

Fig. 13  Intraoperative view following primary neurorrhaphy of transected facial nerve (left) and re-anastomosis of transected parotid duct (right) over a stent due to a penetrating cheek injury.

The risk of sialocele formation. 26 Injection of botulinum toxin A into the gland (50–100 units) has proven to be more efficient and less uncomfortable in minimizing saliva secretion compared with restriction of oral intake or anti-sialogogue use. 27 In cases of extensive glandular damage, subsequent surrounding edema may result in obstruction of the PD and possible sialadenitis. Therefore, the PD should be cannulated to maintain the patency of the lumen in these situations. 26 It is advised that the canal be left in for 2 weeks in this scenario. 28

Parotid Duct Injury

An initial attempt at diagnosing injury to the PD may be performed by massaging the parotid gland. Subsequent pooling of saliva in the surgical field is indicative of damage to the ductal structure. Alternatively, cannulating the distal opening of the duct with a pediatric catheter or lacrimal probe followed by injection of saline may indicate ductal injury by the presence of saline from the distal severed duct within the surgical field. 23 If there is no evidence of ductal compromise, it is recommended that the distal catheter be left in place for 1 week to prevent obstruction. 26, 28 A flexible sialendoscope may also be employed to locate the proximal and distal portions of the transected duct and allow for direct visualization of the ductal system. In cases of repair via re-anastomosis, sialendoscopy is useful for assessment of the anastomosis both peri and postoperatively. 29

When both stumps of the injured PD can be identified and the defect is less than 10 mm, primary re-anastomosis is the preferred method of repair.26, 27 Suturing over a cannula, introduced through the intraoral orifice, is thought to reduce the incidence of ductal stenosis. Venous, urethral, and epidermal catheters have been used as stents. 30–32 Öztürk et al described using an intubation tube cuff as a cannula. 33 Silk or nylon splints have also been used to maintain duct integrity during suturing. 27, 34

In our practice, we prefer the Walvekar Salivary stent (Hood Laboratories, Pembroke, MA), which is specifically designed and anatomically suited for use within the PD. It comes in sizes of 1.0, 1.2, and 1.4-mm diameter, with the largest having the ability to be used as an irrigating catheter due to its luer-lock mechanism. Its defining feature is its dual flanges that are oriented for fixation to the floor of mouth (for submandibular duct cannulation) on one end and the buccal mucosa (for PD cannulation) on the other, preventing migration (► Fig. 14). Each stent comes with a calibrated guide wire that allows sizing of the length of the stent (► Fig. 15). After insertion of the stent to span the transection, anastomosis over the stent may be performed with 7–0 or 8–0 nylon (► Fig. 16). It is secured to the mucosa using a loop stitch with a 4–0 nylon suture on a small cutting needle (► Fig. 17). If using a stent for repair, it should be left in place for a minimum of 2 weeks to allow for healing, though we recommend leaving it in place for 4 to 6 weeks when placed for traumatic etiologies. If the ductal defect is greater than 1 cm, reconstruction with an autologous facial vein graft has been performed successfully and may be considered. 26, 35, 36

Damage to the distal portion of the duct (region C) may make identification of the severed ends of the duct more difficult and primary re-anastomosis may not be feasible. In this case, parotid diversion can be performed by transporting the proximal ductal stump through the buccinator muscle. The distal portion of the duct is then marsupialized to the buccal mucosa using 8–0 nylon suture, creating an artificial opening in the oral mucosa anterior to the natural papilla of the PD. 26, 37 It is advisable to leave a stent within the new orifice for 2 weeks to prevent stomal stenosis or obliteration. 38

If there is extensive ductal trauma, making re-anastomosis or oral re-implantation impossible, the proximal end of the duct should be ligated or clamped. The resultant obstruction causes increased pressure within the parotid parenchyma, culminating in glandular atrophy. This process reduces the risk of sialocele and fistula formation. 20, 23 Initially, swelling of the gland can be painful. Application of a pressure dressing can help promote atrophy. 20
Complications

Stenosis
Post-surgical stenosis of the PD, resulting in obstruction, often presents as recurrent painful swelling of the parotid gland following food intake. Most stenotic lesions occur in the proximal or middle third of the duct. In the past, conventional sialography has been used for diagnosis, but has mostly been abandoned due its need to be used with radiographic contrast. Ultrasound is a first-line diagnostic tool if stenosis is suspected. Alternatively, magnetic resonance sialography may be used. Sialendoscopy is the gold standard to establish final diagnosis and characterize the stenosis to plan accurate treatment. Depending on the severity of the stenosis, the current prevailing method of treatment involves a combination of ductal irrigation with a steroid solution (e.g., 16 mg dexamethasone in isotonic saline) in conjunction with sialendoscopic-controlled dilation. For persistent stenosis, balloon dilation should be attempted. Ultimately, for high-grade stenosis, a stent may be kept in place for 4 to 6 weeks.

Sialoceles and Fistulas
A sialocele results from the accumulation of saliva in the tissues surrounding the parotid gland. It typically presents as a soft, tender, mobile mass in the region of the mandibular ramus approximately 8 to 14 days after initial injury. While sialoceles are often mistaken for hematomas, an aspirate with an amylase concentration above 10,000 U/L can confirm the diagnosis. Treatment depends on the time of sialocele presentation. In the case of an acute process, the wound should be surgically explored. Any damage to the parotid gland that is encountered should be cleaned and the parotid fascia should be sutured. If a lacerated duct is found, reapproximation is indicated. If unsuccessful, the proximal portion of the duct should be ligated and the sialocele should be drained intraorally via catheter.

Treatment of a late-forming sialocele consists of repeated percutaneous aspirations and pressure dressing application to induce glandular atrophy. Anti-sialagogues, such as glycopyrrolate (2 mg twice daily) or scopolamine (1.5 mg transdermal patch every 48 hours), may be used as adjunctive therapy. In addition, antibiotics that cover Staph
Aureus, H. Influenzae, and gram-negative anaerobes may be considered (e.g., Augmentin 850 mg BID). Some authors recommend parenteral nutrition to reduce salivary and allow for more rapid sialolecue resolution. A fistula is a communication between the skin and the parotid gland that usually develops within the first week of injury. Sometimes a fistula may be due to a long-standing sialolecue that ruptures the skin. Patients often notice clear secretions with milking of the gland and with mastication. If a fistula is an immediate complication, surgical re-exploration should be performed. Closure of the parotid capsule is indicated in the case of parotid gland laceration and primary re-anastomosis is preferred if ductal injury is encountered. The proximal portion of the duct should be ligated and secretions drained intraorally via catheter if re-anastomosis is not feasible. Treatment of a persistent fistula involves pressure dressings and anticholinergic medications.

Several studies have shown intraglandular injections of botulinum toxin type A to be effective in the treatment of both sialolecues and fistulas. Lovato et al described a technique used to achieve effective healing of post-parotidectomy fistula in which 10 to 20 units of botulinum toxin, fractionated into three doses, were injected transcutaneously into the parotid gland (two for the superior and one for the inferior lobe). Injections may be performed in conjunction with electromyographic monitoring to prevent undesirable weakening of the masseter and pterygoid muscles. Alternatively, ultrasound may be employed for guidance. Effects take place in 2 weeks and last between 2 to 3 months.

If initial surgical and pharmacologic measures fail to resolve a sialolecue or fistula, more aggressive interventions include radiation therapy or parotidectomy. Low-dose radiation acts to reduce glandular secretions by inducing fibrosis, though it has largely been abandoned due to the risk of malignancy. Tympanic neuromyctosis, which involves the transection of Jacobson’s nerve, a branch of the glossopharyngeal nerve, acts to reduce parasympathetic innervation to the parotid gland. Ultimately, superficial or total parotidectomy may be needed for refractory cases.

Conclusion

Acute soft tissue trauma to the head and neck is among one of the most common reasons for emergency department presentation. Due to the risk of significant long-term functional and cosmetic compromise, evaluation of a patient who has suffered facial trauma should always include an FN exam and carry a low threshold of suspicion for PD injury. In FN repair, neurorrhaphy technique is primarily based on the ability to obtain tension-free anastomosis and outcomes are principally related to the timing of repair. PD injuries are generally repaired based on the site of ducal injury. Therefore, a fundamental knowledge of the anatomy of FN and PD and symptoms of their injuries is paramount in delivering optimal surgical results.

Conflict of Interest
None declared.

References


