An Update on Midface Advancement Using Le Fort II and III Distraction Osteogenesis

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Background

In 1901, Rene Le Fort published a report on fractures of the craniofacial skeleton.1 He defined and differentiated fractures patterns of the adult craniofacial skeleton, developing the now commonly used terminology Le Fort I, II, and III.2 Gilles, Tessier, and others applied Le Fort’s principles to facial osteotomies, and today they are employed to treat patients with craniofacial syndromes such as Crouzon, Apert, or Pfeiffer syndrome who present with midfacial retrusion, shallow orbits, exorbitism, malocclusion, obstructive sleep apnea and facial imbalance.2–8

Classically, the Le Fort II osteotomy involves pyramidal-shaped osteotomies. The bony cuts traverse the nasofrontal suture, the nasal process of the maxilla, the medial orbital wall at the lacrimal bone, the orbital floor, inferior orbital rim and the zygomaticomaxillary suture (► Fig. 1). Both pterygoid plates also need to be osteotomized posteriorly. Le Fort III osteotomy, also known as craniofacial dysjunction, can be performed via an intracranial or extracranial approach. The osteotomy traverses the nasofrontal suture, through the fontal process of the maxilla down the medial orbital, through the orbital floor, across the zygomatic frontooral suture and zygomatic arch (► Fig. 2). Similarly, both pterygoid plates need to be osteotomized posteriorly.4,6

Concepts of Midfacial Distraction

Introduced in 1993 by Cohen et al,9 midfacial distraction osteogenesis (DO) has since become a powerful tool in the armamentarium of the craniomaxillofacial surgeon. Over the past two decades, DO has become an adjunct technique to conventional advancement procedures associated with grafting and rigid fixation.10 The main advantages of midfacial DO over traditional osteotomies include the possibility of producing larger movements, as well as a decreased infectious risk and relapse rate due to the progressive distraction and formation of bone. Distraction osteogenesis also provides for gradual expansion of soft tissues, often resulting in pleasing repositioning of the lips, cheek mounds, and eyelids. Moreover, less subperiosteal dissection is performed, operative procedures are shorter in duration, intraoperative blood loss is reduced, the potential use of bone grafts for stabilization purposes is eliminated, while at the same time eliminating potential donor site morbidity. Hospital stays have also been reported to be shorter.11–19

The general rule of thumb is that conventional advancement of the midface is indicated in patients requiring less than 8 to 10 mm of advancement. Midface DO is often indicated in patients with craniofacial syndromes or any patients with severe midfacial retrusion requiring advancement of more than 10 mm.15,19 Several reports in the literature demonstrate that DO can achieve advancements exceeding the advancement of the traditional procedure by two- to threefold.11–13,20–23 That stated, some feel that DO is indicated because of its positive soft tissue effects, even in smaller advancements.
Indications

Le Fort II
Midface advancement using Le Fort II DO is commonly indicated in patients with midface hypoplasia involving the maxilla and nasal complex, with an acceptable position of the zygomatic complex and orbits. Those patients often will have an anterior cross bite, class III malocclusion, and obstructive sleep apnea. The aims of the Le Fort II DO are to re-establish ideal overjet and overbite and achieve an acceptable occlusion with good functional class I occlusion. At the same time, the Le Fort II allows for lengthening of the nose, a pleasing effect especially in syndromic patients with a classic “dish face” and short nose such as Apert syndrome.

Le Fort III
Le Fort III DO is indicated in syndromic craniofacial patients with midface hypoplasia involving the nasal, maxillary and zygomatic complex, shallow orbits, exophthalmos, upper airway obstruction and obstructive sleep apnea, class III malocclusion and an overall severe facial aesthetic imbalance. Advancement of the midface with Le Fort III expands the nasopharynx and oropharynx, often allowing for tracheostomy decannulation. In addition, by advancing the midface using Le Fort III DO, the abnormal proptotic position of the globe relative to the orbital rim will be corrected preventing amblyopia, corneal exposure with subsequent exposure keratitis, keratoconjunctivitis sicca and infection leading to corneal ulceration, cataracts, and possibly vision loss.

Often a Le Fort III is converted to a monobloc procedure when the retrusive midface is also associated with a retrusive forehead. In Fig. 3, we present a 12-year-old girl with Crouzon syndrome and associated midface hypoplasia involving the nasal, maxillary and zygomatic complex, who underwent successfully a subcranial Le Fort III with DO.

Designer Osteotomies
In rare instances, the traditional Le Fort osteotomies do not ideally treat all patients with craniofacial syndromes. The concept of designer osteotomies can be applied more broadly to include nonclassic midfacial osteotomies in syndromic patients. The following are two examples.

Le Fort II with Zygomatic Repositioning
Hopper has extensively studied the unique morphology of the Apert craniofacial skeleton, reporting a “dual axial and sagittal plane concavity that creates an abnormal face in an abnormal position.” In this subset of patients, a simple Le Fort III does not address the axial concavity of the maxilla. His approach consists of a Le Fort III osteotomy and acute repositioning of the zygomas (osteotomies illustrated in Fig. 4), which are fixed laterally with titanium plates to the lateral orbital wall and frontal bone, as well as a Le Fort II osteotomy segment that is externally distracted using a custom maxillary splint. Using this combined Le Fort II distraction and zygoma repositioning, the orbital–malar relationship is preserved, no iatrogenic periorbital deformity is created, the Le Fort II segment is distracted significantly more than the zygoma, the midface is lengthened, and the anterior open bite is closed.

Combined Monobloc and Le Fort II
Along the same lines, Taylor and colleagues described a different technique to address the facial dysmorphism of Apert patients, who require, as we mentioned above, differential advancement of the forehead, orbits, and midface to correct fronto-orbital retrusion as well as to lengthen the midface and close the anterior open bite. This technique involves a monobloc minus Le Fort II to address these differential movements in a single operation. Advancement
of the forehead and midface is accomplished with the monobloc, while the nose is lengthened and maxilla rotated using the Le Fort II distraction. Similar to Hopper's approach, the monobloc Le Fort II adapts classic craniofacial osteotomies to fit the unique morphology of a given situation. In Fig. 5, we present an 11-year-old girl with Apert syndrome who presented to our clinic with the typical the Apert phenotype, including bicoronal synostosis, papilledema, exorbitism, a severely retruded midface, a shortened nose, and a severe class III malocclusion. To correct her turribrachycephaly and exorbitism, a monobloc distraction with a horizontally oriented vector was planned with internal distractors. To
midface retraction and ocular proptosis/exophthalmos may benefit from a relatively early procedure. On the other hand, patients with less severe morphology and symptoms may benefit from these procedures at skeletal maturity. Each patient should be approached in an individual manner.

External versus Internal Distractors for Le Fort II and III

Distractors come in various shapes and forms, but they can generally be classified as either internal distractors or external distractors (halo-like devices). The advantages and disadvantages of each are discussed in the following paragraphs.27

The main advantage of an internal distractor is that it is less conspicuous and impacts less the patient’s daily activities. These devices are relatively small and less intrusive for patients and families. Major disadvantages include uniplanar distraction vector, inability to manipulate the distraction vector postoperatively, and a slightly increased infection rate.38 Internal devices require a second procedure under anesthesia for removal, which can also be challenging.27 Various groups36,37 have examined the role of resorbable internal devices, however, no comparative studies have been performed and no long-term follow-up studies have been published.

In 1997, Polley38 was the first to describe the use of an external “halo-type” distractor for Le Fort III distraction. Although rigid, external devices allow for easy adjustment in the postoperative period, often in more than one vector. This allows for “orthodontic” adjustment of the distracted segment in multiple vectors to maximize its final position. Additionally, halo-type distractors provide a central “pull” rather than a peripheral “push,” which serves to further unfurl facial concavity often present in syndromic patients. The ability to minimize buried hardware, especially in the region of the bony regenerate, helps to minimize infectious complications as well as maximize bone formation. Generally, they are easy to apply and easy to remove. Disadvantages include the psychosocial discomfort of wearing a large external device, risk of accidental dislodgement, possible infections around the pin sites, scars in the scalp, and the risk of penetration transcranially of the fixation pins.27

Two studies, by Gosain et al39 and Fearon,18 compared outcomes between the use of an external and internal devices. Gosain suggests the use of an external device in older patients and found similar outcomes between the two devices.39 Fearon reported superior results with external distractors compared with internal ones when performing Le Fort III distraction.18 At the Children’s Hospital of Philadelphia, we have noticed a trend toward using external devices, likely for all of the reasons noted above.

Author’s Preferred Technique

Le Fort II Distraction Osteogenesis

Le Fort II osteotomies and DO is performed under general anesthesia via a standard zig-zag coronal incision for exposing...
The nasion and medial orbital region. The maxillary antrum and inferior portion of the zygomatic body are exposed via an upper gingivobuccal incision. The upper gingivobuccal sulcus incision is made along the maxillary vestibule from first molar to first molar. Once the nasion, maxillary antrum, and zygomaticomaxillary buttresses are well exposed, reciprocating standard osteotomies following Le Fort II design are performed. The osteotomy through the nasion is made through the coronal incision, slightly below the nasofrontal suture. The nasal septum is cut from above with a guarded osteotome. Osteotomies of the frontal process of the maxilla, the medial orbital wall at the lacrimal bone, the orbital floor, inferior orbital rim, and the zygomaticomaxillary suture are then performed through the gingivobuccal incision. Both pterygoid plates are osteotomized posteriorly using a curved osteotome. Rowe-Kiley disimpaction forceps are then used to gently verify complete mobilization of the Le Fort II segment. Once complete mobilization is verified, an external distractor is applied, with two fixations pins that are applied bilaterally paranasally at the pyriform aperture. For additional control of the distraction segment, a fixation pin can be added at the region of the nasion. Following abundant irrigation, layered closure of the coronal incision and gingivobuccal sulcus incision is performed. Distraction is begun after a 5-day latency period and proceeds at a rate of 1 mm per day. Our consolidation period lasts approximately 2 to 3 months and the external distraction device is then removed in the operating room. Close consultation with a craniofacial orthodontist can help guide the occlusion either with Tads or guiding elastics.

**Le Fort III Distraction Osteogenesis**

Subcranial Le Fort III DO is performed under general anesthesia via a standard zig-zag coronal incision for exposure of the lateral frontotemporal skull, nasion, temporal fossa, lateral orbital region, zygomatic arch, and zygomatic body. Using a reciprocating saw as well as an ultrasonic saw, standard osteotomies are performed through the zygomatic arch, frontozygomatic suture, floor of the orbit, and nasion. Care is taken to avoid transcranial migration of the saw during this...
portion of the osteotomy. In the midline, the vomer and ethmoid are separated from the cranial base using a guarded swallow-tail osteotome. Both pterygoid plates are osteotomized posteriorly via transmucosal approach or the coronal approach. Rowe-Kiley disimpaction forceps are used to gently verify complete mobilization of the Le Fort III segment. Once complete mobilization is verified and prior to the application of the external distractor, a titanium mesh may be applied to both temporal fossas, to prevent transcranial migration of the distractor fixation pins (in patients with thin bone or cranial defects in the temporal region). Two fixation pins are applied bilaterally paranasally at the pyriform aperture; for added control, additional fixation pins can be applied in the region of the nasion. Those pins will act as bony anchors for the distraction device. Following abundant irrigation, layered closure of the coronal incision is performed and the external distractor is applied.

Complications

Complications Related to Le Fort II and III Osteotomies

Complications with Le Fort II and III osteotomies with subsequent DO are similar to the ones associated with traditional Le Fort II and III,28,40–43 and include infraorbital nerve injury, orbital/globe injury, excessive bleeding, and infection. Dural injury with subsequent cerebrospinal fluid leak is rare but can occur. Other complications include injury to the infraorbital nerve, orbital/globe injury, strabismus, partial anosmia, and localized infections.

Complications Related to Distraction Osteogenesis

Despite technique and device refinements, midfacial DO remains associated with high complication rates, up to 30 to 40%.16,44,45 These complications range from superficial surgical site infections, mostly around the pin sites, to deep soft tissue infection and abscess formation. Specific to the devices, mechanical failure, pin loosening, frame migrations, and finally intracranial fixation pin migration with dural violation, in the setting of halotype distractor can occur (►Fig. 6A).44,46,47 Management of this latter complication by the application of temporary titanium meshes bilaterally in both temporal areas hosting the fixation pins is demonstrated in ►Figs. 6B and 6C.

Other complications associated with distraction include asymmetric advancement and severe infections requiring surgical drainage and hospitalization. Despite these complications, while comparing midfacial distraction to traditional advancement, Fearon13,14 reported lower complications rates as well as shorter hospital stay in the distraction group.

Authors’ Experience and Complication Rate

From 1999 to 2012, 54 patients underwent midface DO at the Children’s Hospital of Philadelphia, including 23 patients with Apert syndrome, 19 with Crouzon syndrome, and 10 with Pfeiffer syndrome, as well as 2 patients with other craniofacial syndromes. Thirty-three patients underwent a total of 34 subcranial Le Fort III distraction procedures and 21 underwent 21 monobloc distraction procedures. The average age at surgery was 8.0 years (range: 4.0–17.7), while average time between distractor placement and removal was 102.9 days. Thirty procedures were performed with external halo-type distractors (18 Le Fort III and 12 monoblocs), while 25 were performed with buried midface distractors (16 Le Fort III and 9 monoblocs). We identified 19 distractor-related complications: There were a total of 10 (18.2%) in the halo group including 5 (9.1%) requiring separate operative intervention, and 9 (16.4%) in the buried distractor group including 6 (10.1%) requiring separate operative intervention. Major infections were more common in the buried distractor group (n = 8) compared with the halo distractor group (n = 3; p = 0.048). There were four (7.3%) patients in the halo group who had malposition or transcranial pin migration related to postoperative positioning or falls that required operative repositioning. Our experience demonstrates that higher rates of halo displacement requiring surgery are offset with lower rates of infections compared with internal devices.

Relapse and Postsurgical Growth

The main advantages of midfacial distraction over traditional osteotomies and advancement include the possibility of producing larger movements, but also a decreased relapse rate due to the progressive distraction and formation of bone and progressive soft tissue expansion.11–13,20–23,48 Since the introduction in 1993 of midfacial DO,9 all reports12,18,35,49–55 are unanimous with regards to the absence or minimal relapse after DO compared with a higher relapse rate in the traditional acute advancement group. The stability of Le Fort II or III DO advancement is well documented in the literature.

Fig. 6 Example of a complication associated with an external halo-type distractor. (A) Axial view of computed tomography scan demonstrating intracranial fixation pin migration with dural violation. (B,C) Management of this latter complication by the application of temporary titanium meshes bilaterally in both temporal areas hosting the fixation pins.
particularly in the long-term outcomes studies published by the group from New York University\textsuperscript{49–51} and the Dallas group.\textsuperscript{18,32,55} However, they demonstrated that although the midface is stable in its new position, it has absent or minimal horizontal and sagittal postsurgical growth and minimal vertical growth. Relapse seen following midfacial distraction is often a misdiagnosis, with “pseudo-relapse” being actually observed. What is possibly seen is absence of growth of the midfacial distracted segment while the mandible is growing normally. This can be easily corrected with a Le Fort I procedure.

Moreover, when considering postsurgical growth of patients after Le Fort II or III, it is important to recognize which patients are evaluated for growth. For instance, one cannot expect a patient with a craniofacial syndrome and severe midface retrusion to develop midface growth following midface DO. Various studies evaluating facial growth in patients with craniofacial syndromes have reported minimal or absent horizontal and sagittal growth, while vertical growth seems to be preserved.\textsuperscript{32–34,49,56–58} Along the same lines, Fearon\textsuperscript{13} concludes that the absence of growth in this specific patient population is intrinsic to the disease process and not to the procedure performed. Thus, with regards to these findings, overcorrection and overdistraction in this subset of patients should be strongly considered. As mentioned previously, timing of the midface procedure should be bound to clear indications. For example, patients with severe obstructive sleep apnea (OSA), midface retrusion, and ocular proptosis benefit from a relatively early procedure, while patients with less severe symptoms and moderate class III malocclusion may benefit from these procedures at skeletal maturity.

**Future Directions**

As with any procedure that involves patient compliance, distraction may benefit from removal of the responsibility of “turning a screw” through the creation of a motor-driven, computer-controlled device. Inclusion of a wireless micro-motor would eliminate concerns about compliance as well as potentially increase the accuracy of distraction. This would likely lead to an improved experience for families.

Improved engineering of devices will help minimize complications and improve outcomes. Smaller, less intrusive, internal devices would be better tolerated by patients. Smaller devices can be worn out of plain sight and impacts minimally the patient’s daily activities. Ideally, these smaller internal devices would also allow for multidirectional adjustment of the distraction vector. Along those lines, resorbable devices have been introduced.\textsuperscript{36,37} CAD/CAM software can also be used for virtual surgical simulation of the procedure. Cutting and positioning guides improve the accuracy of the osteotomies and distractor placement, potentially improving the final aesthetic outcome of the reconstruction.\textsuperscript{59,60}

Enhancement of bone biology—through growth factor manipulation, stem cell enrichment, or genetic manipulation—may shorten treatment regimens. Shortening the consolidation period would allow for an earlier removal of the distractors, allowing patients to return to their regular activities sooner. Significant research is currently underway evaluating various growth factors that will allow for a faster bony formation and consolidation.\textsuperscript{61–66}

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