In-office use of a steroid-eluting implant for maintenance of frontal ostial patency after revision sinus surgery

Agnieszka Janisiewicz, M.D.,1 and Jivianne T. Lee, M.D.1,2

ABSTRACT

Achieving long-term, successful outcomes with endoscopic sinus surgery (ESS) can be challenging in patients with recalcitrant chronic rhinosinusitis (CRS). Local complications, including scar formation and ostial stenosis, can lead to recurrent blockage and subsequent relapse. The frontal sinus is particularly vulnerable to surgical failure given its narrow outflow and inaccessibility to topical therapies. The advent of steroid-eluting sinus implants has enhanced ESS outcomes, with significant reductions in synechiae, inflammation, and secondary postoperative interventions when placed in the ethmoid cavity. However, use of this technology in the frontal sinus has yet to be described. The purpose of this report is to present two cases, in which in-office frontal placement of a mometasone furoate (MF)-eluting implant facilitated maintenance of ostial patency after revision ESS. The clinical presentation, in-office intervention, and treatment outcomes were examined. Two patients (male, 63 and 68 years of age) with a history of multiple ESS presented with recurrent unilateral frontal headache refractory to medical therapy. Nasal endoscopy/imaging revealed frontal sinus outflow obstruction. Both declined revision ESS under general anesthesia and underwent endoscopic frontal sinustomy/ostial dilation in the clinic. A MF-eluting implant was placed in the frontal sinus at the end of the procedure, with preservation of ostial patency upon last follow-up at 3 and 11 months, respectively. In-office placement of a MF-eluting implant successfully maintained frontal ostial patency in patients with a history of multiple ESS. Additional randomized trials are necessary to determine statistical significance, cost-effectiveness analysis, and long-term efficacy of frontal sinus implantation.

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Chronic rhinosinusitis (CRS) represents one of the most common health care problems in the United States, with significant loss of productivity and quality of life associated with the disease.1,2 In recent years, much knowledge has been gained regarding the multifactorial etiology of CRS, with both host (i.e., allergy, immunodeficiency, ciliary immotility, anatomic obstruction, etc.) and environmental factors (i.e., bacterial/viral infections, biofilms, and pollutants) believed to contribute to its pathogenesis. Current treatment of CRS has focused on addressing such underlying issues as well as attenuating inflammation. To that end, a broad spectrum of therapeutic agents have been used in the medical management of CRS, including antibiotics, corticosteroids, antihistamines, saline irrigations, immunomodulators, etc.3,4 When such medical therapy has failed, endoscopic sinus surgery (ESS) has been employed as an effective second-line intervention, with reported success rates of 76%–98%.5 Focused removal of diseased tissue and judicious widening of natural drainage pathways during ESS help to relieve sinonasal obstruction, reestablish ventilation, and facilitate mucociliary flow.

However, achieving long-term, positive outcomes with ESS is contingent upon optimization of the wound healing environment during the postoperative period.6 Local complications, such as scar formation, middle turbinate lateralization, and stenosis of surgically enlarged ostia, can lead to recurrent blockage and eventual surgical failure. Residual inflammation can also impede mucosal recovery and incite polypoid disease, further compromising surgical results.6 A myriad of postoperative strategies, both mechanical and pharmaceutical, have been developed to mitigate such issues, including stents/spacers, steroids, and medicated irrigations.7

In 2011 a steroid-eluting sinus implant was developed (PROPEL, Intersect ENT, Palo Alto, CA) and approved by the United States Food and Drug Administration (FDA) for use in the post-ESS ethmoid cavity.8 The implant is comprised of a biodegradable polylactide-co-glycolide polymer matrix infused with 370 μg of mometasone furoate (MF), which is gradually released to surrounding tissues over a 30-day period. Such technology has enabled targeted steroid delivery directly to sinus mucosa with simultaneous stenting of the sinonasal cavity. This dual functionality has led to significantly enhanced ESS outcomes with implant use,
as demonstrated in three clinical trials and a meta-
analysis of the aggregate data.8–14 Significant reduc-
tions in adhesion formation, middle turbinate lateral-
ization, and polypoid change as well as the need for
secondary postoperative medical (oral steroid admin-
istration) and surgical interventions (lysis of adhe-
sions) have been reported with ethmoid implanta-
tion.8–14

In 2012, the FDA approved a smaller version of the
ethmoid implant which contained the same dosage of
MF as the original product but featured a reduced
scaffold size (PROPEL mini, Intersect ENT, Palo Alto,
CA). Although designed for deployment in a narrower
ethmoid cavity, its diminished profile has made it ame-
nable to off-label application in the frontal sinus. How-
ever, use of the steroid-releasing implant for the frontal
sinus has yet to be described. The purpose of this
report is to present two cases in which in-office place-
ment of a MF-eluting implant successfully maintained
frontal ostial patency in patients with a history of re-
vision ESS.

CASE REPORT

A 63-year-old male presented to our clinic with left-
sided frontal pressure and headache for six months. He
also complained of intermittent purulent rhinorrhea
from the left, but denied having any nasal obstruction,
nasal congestion, postnasal drip, or visual changes. His
previous surgical history was significant for three
Lynch procedures (two on the left, one on the right)
from 1992 to 1993 as well as a left revision ESS in 1995
for CRS. Since the most recent procedure, the patient
noted longstanding anosmia but otherwise had no
complaints until the time of presentation.

On nasal endoscopy, the patient was found to have a
stenosed antrostomy with a residual uncinate process.
There was also evidence of a partial ethmoidectomy
with remaining partitions along the anterior skull base.
The middle turbinate appeared to have been previ-
ously resected. No frontal sinus outflow tract could be
visualized, and thick synechiae could be seen obstruct-
ing the frontal recess. After a three-week course of
broad spectrum antibiotics, a computed tomography
(CT) scan was obtained. Complete opacification of the
left frontal sinus was shown with an absent lateral
wall, corresponding to the superomedial aspect of the
orbit (Fig. 1, A–C). Due to his persistent symptoms and
radiographic findings, the patient elected to proceed
with left revision endoscopic sinus surgery. Triplanar
stereotactic imaging was performed in preparation for
computer-assisted surgical navigation.

Intraoperatively, significant neoosteogenesis was en-
countered within the frontal recess, requiring extensive
drilling to open the frontal sinus outflow tract (Fig. 2
A). Once all the bone was removed, copious mucopu-
rulent secretions began to drain from the frontal sinus
proper (Fig. 2 B). Specimens were sent for aerobic and
anaerobic cultures which grew *Staphylococcus aureus*. A
Draf 2B frontal sinusotomy was then completed, with a
70° diamond burr and angled instrumentation used to
take down the nasofrontal beak and enlarge the frontal
sinus ostium (Fig. 2, C and D). At the conclusion of the
procedure, inspection with a 70° 4-mm telescope re-
vealed a patent frontal sinus neostium (Fig. 2 E). No
stent nor packing materials were placed. There were no
complications, and the patient was discharged home
with a steroid taper and a three-week course of antibi-
otics.

However, the patient missed all of his subsequent
follow-up visits for postoperative debridement. He
then returned to our clinic two months later, complain-
ing of recurrent left-sided frontal pressure and discol-
ored rhinorrhea. On nasal endoscopy, mucopurulent
drainage was present, and the frontal sinus outflow

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**Figure 1.** Preoperative coronal (A and B) and axial (C) CT images demonstrated complete opacification of the left frontal sinus (start) with an absent lateral wall, corresponding to the superomedial aspect of the orbit (arrows).
tract could no longer be visualized (Fig. 3 A). Cultures were obtained and grew *Streptococcus pneumoniae* and *Pantoea* species. Under local anesthesia, frontal sinus instruments as well as balloon dilation were used to reopen the ostium in the clinic. This was repeated on multiple appointments over the span of six to eight weeks and the patient placed on culture-directed antibiotics. Nevertheless, the ostium continued to close in between visits, leading to persistent infection. Ultimately, due to issues with patient compliance, a silastic stent was placed to maintain frontal sinus ostial patency (Fig. 3 B).

The patient did well for eight months, while continuing saline and budesonide rinses, but then returned once again presenting with left-sided forehead/periorbital pressure and discolored rhinorrhea. Inspection with a 70° endoscope revealed the stent to still be in the proper position with a stenosed but patent frontal ostium. However, the stent was clearly infected with yellow, mucopurulent secretions surrounding the foreign body (Fig. 4 A). The stent was removed, revealing inflamed mucosa lining the frontal sinus (Fig. 4 B). At this point, the patient was offered a Draf III procedure, but he declined to pursue any further surgical intervention. The pus was suctioned and sent for culture, which grew *Pseudomonas putida*, *Stenotrophomonas maltophilia*, and *S. aureus*. The frontal sinus ostium was enlarged using balloon dilation. A steroid-eluting sinus
implant (PROPEL mini, Intersect ENT) was then deployed into the frontal sinus (Fig. 5 A). The implant was positioned such that its inferior edge abutted the frontal ostium circumferentially (Fig. 5 B), and the remainder interfaced with the inflamed frontal sinus mucosa to optimize drug delivery (Fig. 5 C).

Three and six weeks after implantation, the frontal ostium remained open with residual fragments still visible (Fig. 6, A and B). By 10 weeks, the implant had resorbed completely (Fig. 6 C), with frontal ostial patency maintained upon last follow-up at 11 months (Fig. 6 D). During the follow-up period, clear mucus would be suctioned at times around the ostium, but no crusting ever developed and no debridements were necessary to keep the stent patent before its absorption. The implant resorbed completely by 10 weeks without the need for removal of fragments. The patient has had no evidence of recurrent infection and has not required any further topical or systemic steroids since sinus implantation.

Case Two

A 68-year-old man presented to our clinic with worsening left-sided frontal pressure and headache for one year. He had been treated with multiple courses of antibiotics, oral steroids, and six months of twice daily budesonide irrigations without any improvement in his symptoms. His previous surgical history was significant for four bilateral endoscopic sinus surgeries in 1995, 1998, 2001, and 2006 for CRS. Since the most recent procedure, he had sustained longstanding anosmia but denied having any purulent rhinorrhea, post-nasal drip, nasal congestion, obstruction, or visual changes.

Nasal endoscopy showed evidence of a previous ethmoidectomy with a clear posterior skull base, but residual partitions could be appreciated within the frontal recess. A sinus CT was obtained, which demonstrated mucosal thickening of the left frontal sinus (Fig. 7 A). Remnants of bone obstructing the frontal sinus outflow tract could also be appreciated on sagittal views (Fig. 7 B). Left revision ESS was recommended, but the patient declined to undergo general anesthesia and instead elected to proceed with left revision frontal sinusotomy in the clinic. After topical and local anesthesia were administered, frontal sinus instruments and balloon dilation were used to remove the remaining bony ledges within the frontal recess.
and enlarge the natural ostium of the frontal sinus (Fig. 8, A–C). Significant polypoid inflammation was encountered within the frontal sinus (Fig. 8 C). Therefore, a steroid-eluting implant (PROPEL mini, Intersect ENT) was placed (Fig. 9).

At three weeks (Fig. 10 A) postimplantation, the inflammation within the frontal sinus had subsided considerably and the ostium remained open. By seven weeks (Fig. 10 B), the implant had dissolved completely with continued preservation of ostial patency.

Figure 6. Endoscopic photograph at three weeks (A) postimplant showed an open frontal ostium with residual fragments still present. By 10 weeks (B), the implant had resorbed completely, with frontal ostial patency maintained upon last follow-up at 11 months (C).

Figure 7. CT images demonstrated mucosal thickening of the left frontal sinus on coronal (A) and sagittal views (B), with residual bony remnants (arrow) obstructing the frontal sinus outflow tract.

Figure 8. (A) Nasal endoscopy showed residual partitions (arrow) within the frontal recess. (B) After partial removal, the frontal sinus outflow tract could be visualized (arrow). (C) Enlargement of the natural ostium of the frontal sinus (arrow) revealed polyps (star) within the sinus cavity which were debrided.
and no evidence of recurrent polypoid inflammation. No debridements were necessary while the stent was in place to maintain ostial patency, and no topical steroid therapy was given postoperatively. The patient also reported complete resolution of his symptoms after sinus implantation, with no further headaches upon last follow-up at six months.

It should be noted that it is unknown whether balloon dilation and topical steroids, without concurrent sinus implantation would have yielded a similar outcome in this case. The decision to place the steroid-eluting implant was based on the significant amount of polypoid inflammation in the frontal sinus in the context of multiple revision ESS, leading us to suspect he would fail such a trial if attempted.

DISCUSSION

Sinus implantation represents a novel mode of topical drug delivery, in which localized, sustained release of corticosteroids is coupled with mechanical force to facilitate healing of the postsurgical cavity and preserve ostial patency. Currently FDA approved for ethmoid use only, the implant is constructed in an open-lattice configuration, with a unique spring-like design that enables it to physically separate sinonasal tissues as it expands to conform to the shape of the dissected sinus. When compressed, the implant has a diameter of 5.2 mm, which widens to 23 mm once deployed. It is typically inserted intraoperatively under endoscopic visualization at the conclusion of ESS, allowing direct, sustained application of MF to the diseased ethmoid mucosa after surgery.8

This dual functionality of mechanical dilation and antiinflammatory therapy in sinus implantation has been shown to safely and effectively improve postoperative ESS outcomes, as demonstrated in three multicenter studies and a metaanalysis of the aggregate data (more than 200 CRS patients).9–12 In the initial pilot study by Murr et al.,12 a double-blinded randomized controlled trial was performed on 38 CRS patients who had the steroid-eluting implant placed in the post-ESS ethmoid cavity on one side and the identical, non-steroid-eluting implant on the other. At the 60-day follow-up, significant reductions in polyp formation, adhesions, and inflammation were observed on the treatment side compared with controls.12 Likewise, in a follow-up study by Forwith et al. (a single cohort, prospective clinical evaluation of 50 CRS patients, ADVANCE), polypoid tissue, scarring, and middle turbinate lateralization were reported in only 10%, 1.1%, and 4.4% of implanted patients, respectively.9 Statistically significant improvement in patient-reported outcomes and quality of life (Sinonasal Outcomes Test-22, Rhinosinusitis Disability Index) were also described with no deleterious ocular sequelae noted. Subsequently, in a larger double-blinded randomized controlled trial of 105 CRS patients (ADVANCE II), a decreased need for secondary postoperative medical (oral steroids) and surgical (lysis of adhesions) interventions was reported in implanted ethmoid sinuses versus controls.10 Most recently, in a metaanalysis of pooled data (143 CRS patients) from the pilot and ADVANCE II trials, independent panel results revealed statistically significant, relative reductions of 46%, 40%, 51%, and 35% in nasal polyposis, oral steroid administration, lysis of adhesions, and overall postsurgical interventions, respectively, compared with controls.11 Only two device-related adverse events have been reported thus far, an exacerbation of sinus pressure secondary to crusting and granulation tissue leading to implant removal.13 Although rare, hypersensitivity reactions to stent components have been previously reported. However, these were primarily in association with retained nonabsorbable stents containing metal struts.15 Collectively, these studies have demonstrated the clinical benefits of localized steroid delivery via sinus implantation during the postoperative period.

However, these trials primarily investigated the effects of ethmoid implant placement intraoperatively during ESS. Although the potential application of this technology in the office has been postulated, only one series has explored sinus implantation in the clinic setting. This was the recently published prospective multicenter study by Lavigne et al., in which 12 patients with a history of previous ESS and recurrent polyposis underwent in-office placement of MF-eluting implants in the ethmoid sinuses.16 After one month, statistically significant reductions in mean en-
doscopic polyp grades and Sinonasal Outcomes Test-22 scores were observed, outcomes which were sustained over the course of six months. Furthermore, 64% patients had improved to such a degree that they were no longer deemed candidates for revision ESS.16

Thus far, sinonasal application of steroid-eluting implants has been limited to the ethmoid cavity. Use of the MF-releasing spring implant within the frontal sinus has yet to be described. When affected by CRS, the frontal sinus can be particularly problematic secondary to its complex anatomy and narrow outflow tract. Frontal ostial restenosis rates of 33%–50% have been reported after ESS, with 12%–30% requiring revision surgery.17,18 Multiple strategies have been devised to address such scarring during the postoperative period, including various stents, spacers, pharmacologic dressings, etc. with mixed results.19,20 In-clinic procedures, such as balloon catheter dilations, have also been used to treat such recalcitrant disease and recurrent frontal stenosis. Although success in thwarting revision surgery has been reported, a fair proportion of (17%) patients require repeated dilations due to restenosis as seen in the first patient.21

To our knowledge, no studies have been published investigating the use of steroid-eluting implants in the frontal sinus. Here, we report on two patients with a history of multiple sinus surgeries, refractory CRS, and frontal stenosis, in which in-office placement of the MF-releasing implant successfully maintained ostial patency. The PROPEL mini has a compressed diameter of 4.0 mm (versus 5.2 mm), which enlarges to 16 mm (versus 23 mm) upon maximal expansion. Although initially created for patients undergoing less extensive sinus surgery or possessing narrower ethmoid anatomy, its diminished scaffold profile and curved delivery system have made it amenable to frontal sinus delivery. As demonstrated in the cases presented, frontal sinus implantation can be performed in the clinic setting under endoscopic visualization with local anesthesia. Similar to previous published results, the first patient with recurrent frontal sinusitis initially required multiple in-office debridements and balloon dilations to maintain ostial patency.21 However, after placement of the MF-releasing implant, both patients experienced complete resolution of their symptoms and required no additional procedures or oral steroid therapy.

These two patients exemplify the benefits of simultaneously addressing the anatomic obstruction and inflammation that are often the hallmark of CRS. This synergistic combination of ostial dilation and sinus implantation may represent a minimally invasive, in-office therapeutic alternative for management of select patients with medically and surgically refractory frontal CRS. As technologies involving impregnated, biodegradable stents evolve, MF-releasing sinus implants may represent just the beginning of an entire new spectrum of devices harboring both drug-eluting and mechanical stenting capabilities. Future implants may be engineered to conform specifically to the maxillary, sphenoid, and frontal sinuses. In addition, the implant polymer matrix could be loaded with different medications, including high-dose antibiotics, other antiinflammatory agents, or various combinations of drugs. This opens the door to a vast array of topical pharmaceutical therapies that could be administered to each of the sinuses for purposes of facilitating postoperative wound healing, treating recurrent infection, or managing persistent inflammation. In a study by Huvenne et al.,22 doxycycline-releasing frontal stents were found to suppress bacterial growth, reduce matrix metalloproteinase-9 levels, and improve postoperative healing of the frontal region. With such technology, the need for systemic medical therapy and revision ESS may be obviated in certain patients. However, limitations of drug-eluting frontal stents must be kept in mind. Long-

Figure 10. Endoscopic photograph at three weeks (A) postimplant showed an open frontal ostium with residual fragments still present. Note the marked decrease in polypoid inflammation within the frontal sinus. By seven weeks (B), the implant had resorbed completely with continued maintenance of frontal ostial patency after three months of follow-up (C).
term efficacy remains unclear, and establishing a patient outflow tract is still needed before stent placement. Hypersensitivity reactions to implant or drug components may also occur. Furthermore, potential adverse sequelae associated with repeated or multiple sinus implantation must also be elucidated.

CONCLUSION

In-office placement of a MF-eluting implant successfully maintained frontal ostial patency in two patients with a history of multiple ESS and refractory frontal sinusitis. Such intervention may emerge as an effective, office-based therapeutic option for this patient population. Additional randomized trials are necessary to determine statistical significance, cost-effectiveness analysis, and long-term efficacy of frontal sinus implantation.

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