Maxillofacial reconstruction with Medpor porous polyethylene implant: a case series study

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Abstract (J Korean Assoc Oral Maxillofac Surg 2018;44:128-135)

Objectives: The role of alloplastic materials in maxillofacial reconstruction is still controversial. Determining the utility of porous, high-density, polyethylene implants as a highly stable and flexible, porous alloplast, with properties such as rapid vascularization and tissue ingrowth, is crucial in cases of maxillofacial deformities and aesthetic surgery.

Materials and Methods: Thirty high-density porous polyethylene implants were implanted in 16 patients that had been referred to a private office over a three-year period. These implants were used for correcting congenital deformities, posttraumatic defects and improving the aesthetic in nasal, paranasal, malar, chin, mandibular angle, body and orbital areas.

Results: The outcomes of the cases in this study showed good aesthetic and functional results. The majority of patients had no signs of discomfort, rejection or exposure. Two implants suffered complications: a complicated malar implant was managed by antibiotic therapy, and an infected mandibular angle implant was removed despite antibiotic therapy.

Conclusion: Based on the results, the Medpor implant seems to be an excellent biomaterial for correcting various facial deformities. Advantages include its versatility and relatively ideal pore size that allows for excellent soft tissue ingrowth and coverage. It is strong, flexible and easy to shape.

Key words: Maxillofacial prosthesis implantation, Reconstructive surgical procedures, Medpor

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is resistant to infection, exposure and contractile forces.\textsuperscript{4,13,14} Medpor possesses the following mechanical properties: (1) it is easily formed, (2) it is strong enough to be used in non-load bearing areas, and (3) it is readily available as a sterile implant in various pre-formed shapes.\textsuperscript{13}

Our report reviews the use of PHDPE implants for correcting deformities in the orbit (floor, medial, and lateral wall), nasal, malar, chin, mandibular body and mandibular angle in 16 patients.

II. Materials and Methods

We used 30 PHDPE implants (Medpor Biomaterial; Porex Surgical, Newnan, GA, USA) for 16 patients with different types of deformities between the years 2010 to 2012 in Qazvin University of Medical Sciences. All patients were informed about the advantages and disadvantages of these implants. Six sites of deformity (malar area, orbital floor, mandibular ramus and body, nasal, paranasal and chin area) were corrected with prefabricated Medpor. All surgeries were performed under general anesthesia and through intra- or

Fig. 1. Application of a “M” design malar implant and a paranasal implant. A, C. Preoperative lateral view. B, D. Postoperative lateral view. E. Preoperative view from above. F. Postoperative from above.

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Fig. 2. Augmentation of malar bones using two “M” design malar implants and addressing the saddle nose with a nasal radix implant. A. Preoperative three-quarter view. B. Postoperative three-quarter view. C. Preoperative lateral view. D. Postoperative lateral view. E. Preoperative frontal view. F. Postoperative frontal view.

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extra-oral approaches based on the case and minimal manipulation. Implants were inserted in the subperiosteal plane and secured with screws. Before insertion, implants were individually contoured with a scalpel based on deformity geometry, and they were immersed in a gentamicin solution in order to minimize risk of infection. After surgery, regular postoperative protocol, including pain relief and intravenous antibiotic therapy (cefazolin 1 g, four times a day), was prescribed. Patients were discharged with oral antibiotics (cephalexin 500 mg, four times a day) for one week, and mouthwash rinse was prescribed for cases with an intraoral approach.

Patients were recalled routinely after 1, 4, and 12 weeks for postoperative follow-up, and long term follow-up data were collected after 5 to 7 years. Patient satisfaction and objective criteria (asymmetry and displacement, postoperative complications and corrective surgery) were evaluated.

1. Presentation of cases

1) Case 1
A 22-year-old woman with malar and paranasal deficiency. A “M” design malar implant and paranasal implant were used and fixed with screws. (Fig. 1)

2) Case 2
A 20-year-old woman with severe saddle nose deformity as well as malar and infraorbital rim depression. Initially, two “M” design malar implants were used to augment the malar area. The nasal dorsum was corrected by nasal radix and columellar strut implants, increasing the nasal projection and elevation. Finally, we reinforced and reconstructed the upper lateral cartilages using a Medpor implant. (Fig. 2)

3) Case 3
A 25-year-old man complaining of diplopia with an orbital blowout fracture and enophthalmous following a road traffic accident. At first, the zygomatico-maxillary complex fracture was approached. Next, diplopia and enophthalmous were addressed using a Medpor channel implant which is thicker posteriorly and thinner anteriorly, causing the globe to move forward. We used an infraorbital rim incision for accessing the orbital floor and the medial and lateral wall of the orbit. Orbital defects in the medial and lateral wall were corrected by a Medpor implant reinforced with titanium mesh. After three months, the patient had no diplopia or enophthalmous. (Fig. 3)

4) Case 4
A 30-year-old woman complaining of malar deficiency and poor facial contour in the mandibular angle and the lower and upper lip area. A “M” design malar implant was used for malar bone augmentation, and autologous fat was injected to refine the angle, ramus, and around the lips. (Fig. 4)

III. Results

A summary of patients is shown in Table 1. This study included 13 females and 3 males, and the mean age was 26.3±4.4 years. There were three indications for the application of PHDPE implants in our clinic: congenital deformity (26.7%), posttraumatic defect (26.7%), and aesthetic (46.7%).
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Table 2

Table 1. Summary of patients

| No. | Age (yr) | Sex | Site of defects | Etiology | Follow-up (yr) | Complication | Corrective surgery | Asymmetry | Patient satisfaction | No. of implants used |
|-----|----------|-----|-----------------|----------|---------------|--------------|--------------------|-----------|----------------------|---------------------|
| 2   | 23 F     | Malar | Aesthetic      | 5        | No            | No           | No                | No        | Yes                  | 1                   |
| 3   | 28 F     | Malar | Aesthetic      | 7        | No            | No           | No                | No        | Yes                  | 2                   |
| 4   | 32 F     | Malar | Aesthetic      | 5        | No            | No           | No                | No        | Yes                  | 2                   |
| 5   | 25 F     | Malar | Congenital     | 5        | No            | No           | No                | No        | Yes                  | 2                   |
| 6   | 32 F     | Malar | Paranasal      | 5        | No            | No           | No                | No        | Yes                  | 2                   |
| 7   | 20 M     | Mandibular body & angle | Trauma | 5 | No | No | No | Yes | 1 |
| 8   | 20 M     | Mandibular body & angle | Aesthetic | 5 | No | No | No | Yes | 2 |
| 9   | 30 M     | Orbit | Aesthetic      | 7        | No            | No           | No                | No        | Yes                  | 3                   |
| 10  | 25 M     | Mandibular body & angle | Aesthetic | 6 | No | No | No | Yes | 2 |
| 11  | 30 M     | Mandibular body & angle | Aesthetic | 6 | Infection | Angle implant removed Incision and drainage | Yes | No | 2 |
| 12  | 26 F     | Mandibular body & angle | Aesthetic | 5 | Displacement & infection | Incision and drainage | Yes | No | 1 |
| 13  | 24 F     | Paranasal | Congenital | 5 | No | No | No | Yes | 1 |
| 14  | 24 M     | Chin | Aesthetic      | 5        | No            | No           | No                | No        | Yes                  | 1                   |
| 15  | 26 M     | Chin | Trauma         | 6        | No            | No           | No                | No        | Yes                  | 2                   |

(F: female, M: male)

Table 2) Implants were used for different sites of the maxillofacial region according to the following frequencies: malar (50.0%), orbital (10.0%), mandibular body and angle (13.3%), nasal (10.0%), paranasal (6.7%), and chin (10.0%). (Table 3) The mean follow-up was 5.4±0.62 years with complication and satisfaction rates of 12.5% and 87.5%, respectively.

In the postoperative period, two patients suffered complications in the malar and mandibular angle areas. We administered an intravenous antibiotic to these patients, and their wounds were incised and drained. This protocol was successful in one patient, but we had to remove the angle implant in the second patient. All other patients were satisfied with their treatment.

Fig. 4. Using “M” design malar implant to augment malar bone combined with fat injections to the mandibular angle, ramus, and both lips. A. Preoperative lateral view. B. Postoperative lateral view. C. Preoperative view from above. D. Postoperative view from above. E. Preoperative frontal view. F. Postoperative frontal view.

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Table 2. Etiology of deformities

| Reason for implant | No. of implants (%) | No. of cases |
|--------------------|---------------------|-------------|
| Posttraumatic      | 8 (26.7)            | 4           |
| Aesthetic          | 14 (46.7)           | 8           |
| Congenital         | 8 (26.7)            | 4           |

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Table 3. Sites and number of implants used in patients

| Region              | No. of patients | No. of implants (%) |
|---------------------|-----------------|---------------------|
| Nasal               | 8               | 15 (50.0)           |
| Orbit               | 1               | 3 (10.0)            |
| Mandibular body and angle | 2    | 4 (13.3)            |
| Nasal               | 1               | 3 (10.0)            |
| Chin                | 2               | 3 (10.0)            |
| Paranasal           | 2               | 2 (6.7)             |

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IV. Discussion

Facial implants are frequently used for aesthetic purposes, correction of congenital deformities and restoring anatomical harmony after trauma. Facial implants became popular around the turn of the century, but Rousssett was using gold implants in the nose as early as 1828. Joseph used ivory inlays for the nose in 1907. In 1896, Israel used tibial bone for nasal reconstruction, and in 1900, cartilage was used for reconstructive purposes by Von Mangold. Brown et al. reported the advantages of silicone implants in 1953, and silicon is one of the most widely used implant materials today.

An ideal alloplastic implant should be inert, non-carcinogenic, non-inflammatory and non-allergenic. In addition, it should resist mechanical strain and be easy to fabricate and shape. An optimal implant should integrate with the surrounding soft tissue, bone and cartilage.

Autologous materials, despite some disadvantages, still remain the gold standard for craniofacial reconstruction. Increased time and complexity of surgery, donor site morbidity, difficulty in graft shaping, warpage and resorption have been referred to as disadvantages of autologous materials.

This report describes our experience with PHDPE implants (Medpor Biomaterial), an alloplastic implant material that may offer advantages when compared with previously used materials.

Medpor is user friendly, and it may easily be fixed for restoration of a three-dimensional structure. From a physical point of view, Medpor is a pure complex, and it is composed of biocompatible material that is strong and does not easily undergo degradation. These qualities, along with an ability to maintain its initial volume, make Medpor a suitable alternative for autogenous graft or other alloplastic materials. By forming interconnecting pores, the Medpor implants are suitable materials for orbital reconstruction. These pores size range from 160 to 368 μm, and more than half of these pores are larger than 150 μm in diameter. Klawitter et al. and Spector et al. stated that pore size larger than 100 μm promote tissue ingrowth. For example, materials such as Polytef (Gore-tex) have pore sizes much less than 100 μm. Such porosity results in the ingrowth of vascularized tissues into the implant, eventually forming a highly stable complex which is resistant to infection and deformation. But, on the other hand, this soft tissue ingrowth can make surgical removal of porous polyethylene extremely complicated.

Extensive vascular ingrowth into the implant transfers cellular production deep into the implant, which can promote infection resistance. However, Romano et al. managed facial fractures in 140 patients with Medpor implants, and infection was observed in only one case. In another study by Ridwan-Pramana et al., they reported 7.2% infection rate in 69 implants that have been used in 40 post trauma patients.

Medpor implants have been used for nasal deformities. Niechajev used 118 implants in 102 patients for nose deformities, chin hypoplasia and malar hypoplasia. Three rhinoplasty cases were infected, two cases were faced with partial extrusion, and two dorsal and two chin implants were trimmed. Mohammadi et al. used PHDPE in open rhinoplasty. Medpor can be used as a dorsal and spreader graft in the correction of severe nose deformity without noticeable complications, such as infection and extrusion. In the current study, we used 3 implants for nasal reconstruction (dorsal, columellar, and upper lateral cartilage implants) without any complications.

Medpor implants have been used for orbital deformities as well. In a study by Rubin et al., only one orbital implant was infected 1-week postoperatively, and other minor complications included under correction, a symptomatically palpable implant, and transient postoperative chemosis. In different studies by Baj et al. and Rapidis and Day, high density polyethylene implants were recommended for reconstruction of the temporal defect after temporalis myofascial flap transposition. It is an easy and safe method with excellent functional and aesthetic results, and the method has a success rate of 90%. Studies by Rubin et al. and Xu et al. have shown
that porous polyethylene sheets are very reliable materials for reconstruction of orbital blow-out fractures, restoration of orbital volume, and treatment and correction of diplopia and enophthalmous.

Xu et al. explained that overcorrection of 1 to 2 mm is necessary during surgery due to soft tissue swelling or atrophy. They observed no sign of infection in their study. Some authors reported risk of patient dissatisfaction because the rigid nature of porous polyethylene makes it difficult to contour. The current use of computer-aided design/computer-aided manufacturing (CAD/CAM) techniques have facilitated the rapid and precise construction of customized implants. Sun et al. showed that customized titanium mesh can accurately correct enophthalmous accompanying orbital fractures. He et al. stated that computer-assisted surgery can improve the treatment outcomes in delayed orbito-zygomatic fracture with enophthalmous.

In two separate studies by Yilmaz et al. and Lin et al., 26 and 21 patients, respectively, with orbital floor fractures were treated by porous polyethylene implants. The patients had problems such as enophthalmous, diplopia, limited extrinsic ocular motility, hypoglobus and impairment of the infraorbital nerve. They concluded that porous polyethylene sheets are safe, reliable and effective implants. In addition, they reported that the sheets could be used for orbital floor fracture reconstruction without donor site morbidity or need for implant fixation. In a study conducted by Lin et al., orbital infection and/or worsening of diplopia weren’t observed in any of the patients studied. In a study by Yilmaz et al., 4 postoperative infections in 4 patients were managed with antibiotics, and ectropion was formed in 2 cases. Cenzi et al. and Yaremchuk, used 285 Medpor implants in 187 patients and 370 implants in 162 patients, respectively. They concluded that porous polyethylene implants have favorable properties for craniofacial skeletal reconstruction. None of their patients developed complications, such as extrusion, migration and infection.

Use of Medpor implants for areas like the ear, nose and maxilla in syndromic patients is associated with a higher risk of implant failure. A study by Gosau et al. showed fibrovascular integration without encapsulation under light microscopy for the Medpor implant. They detected giant cells on the surface of the implants and evidence of implant material resorption.

An important disadvantage of Medpor implants is their invisibility in radiographic studies, because the Medpor implant shows no contrast. Menderes et al. placed 83 implants in 71 patients for craniofacial reconstruction. Their study showed an increased risk of early and late exposure if the Medpor implant was placed directly under the skin instead of the subperiosteal plane. They preferred autogenous grafts, instead of alloplastic materials, for reconstruction of the nasal dorsum and microtia. Romo et al. stated that if autogenous materials are inadequate or undesirable, the surgeon could use the Medpor implant to reconstruct and support the external nasal valve. On the other hand, Emsen supported the use of E-M shaped septal encircling with Medpor grafts as a safe, effective, reliable and permanent method for crooked nose reconstruction.

In this study, 16 patients with 30 Medpor implants were evaluated. We achieved good aesthetic and facial contour in all patients treated with PHDPE implants. None of the patients developed implant exposure. Two cases of postoperative infection in the malar and mandibular angle area were managed by incision and drainage and antibiotic therapy. Only the mandibular angle implant was removed due to unsuccessful medical treatment.

Based on the findings of this study, we can suggest PHDPE implants, having a low incidence of infection and acceptable aesthetic and functional outcomes.

V. Conclusion

The PHDPE alloplastic implant has a low incidence of infection and excellent cosmetic and functional results, and the implant is an acceptable alternative to existing alloplastic materials.

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Authors’ Contributions

M.K. organized the survey, designed the study, and treated the patients together with P.J. and F.R. M.K. and P.J. revised the manuscript. All authors read and approved the final manuscript.

Ethics Approval and Consent to Participate

This study was approved by the Qazvin University of Medi-
Cal Sciences Ethics Committee (no. IR.QUMS.REC.1394.809). There is no conflict with ethical considerations.

Consent for Publishing Photographs

Written informed consent was obtained from all patients for publication of this article and accompanying images.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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