Prevalence of complications associated with polymer-based alloplastic materials in nasal dorsal augmentation: a systematic review and meta-analysis

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Abstract

Background: Various techniques with different grafts and implants have been proposed to establish a smooth and symmetric nasal dorsum with adequate function. Broadly, two categories of materials have been used in this regard: alloplastic implant materials and autograft materials. The aim of these meta-analyses is to explore the incidence of complications after dorsum augmentation surgery using alloplastic materials.

Materials and methods: After duplication removal 491 papers remained that title and abstract were assessed for eligibility. Regarding the study type, 27 observational studies were included, 21 retrospective and 6 prospective case series. A total of 3803 cases were enrolled in this systematic review and meta-analysis.

Result: Twenty-seven articles reported on complications and outcomes of dorsal augmentation rhinoplasty with synthetic materials. In a random-effects model, the weighted mean percentage was 2.75% (95% CI 1.61 to 4.17%). The weighted mean percentage were 1.91% (95% CI 0.77 to 3.54%), 0.72% (95% CI 0.316 to 1.31%), and 0.78% (95% CI 0.43 to 1.24%) respectively.

Conclusion: The widely used alloplasts were expanded polytetrafluoroethylene (ePTFE), high-density polyethylene, and silicone. The total rates for complications, infection, deviation, irregularity, hematoma, extrusion, and overcorrection were 2.75%, 1.91%, 0.72%, 0.70%, 0.78%, and 0.49%, respectively. The revision rate, based on the random effects model, was 6.40% with 95%CI (3.84 to 9.57).

Trial registration: This meta-analysis was registered at the International Prospective Register of Systematic Reviews (PROSPERO, registration number CRD42020209644).

Keywords: Rhinoplasty, Augmentation rhinoplasty, Alloplastic, Complications, Revision rate of rhinoplasty

Background

Over the time, the different approaches on rhinoplasty have shifted from reductive towards augmentative. The nasal dorsum height and shape, and its harmonious alignment with tip of nose, play a key role in creating perfect esthetic results [1, 2]. In cases with indistinct nasal bridges, dorsal deficiencies, and under-projected nasal dorsum, dorsal augmentation is the recommended procedure [3]. Various techniques with different grafts and implants have been proposed to establish a smooth and symmetric nasal dorsum with adequate function. Broadly, two categories of materials have been used in this regard: alloplastic implant materials and autograft materials [4].

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First is widely used in west while the latter is the preferred item among Asian surgeons [5, 6]. There remains a controversy regarding the selection of the appropriate material with more advantages and lower complication rates. The autologous materials are preferred for dorsal augmentation due to low infection and extrusion rates and high biocompatibility. Although there remains concerns of complications such as major resorption and graft harvesting site morbidity with autologous grafting. Alloplastic materials such as silicone,

**Table 1** Search strategy

| Search criteria                                                                                     |
|-----------------------------------------------------------------------------------------------|
| PubMed (365) \(((\text{alloplast Title/Abstract}) \text{ OR silicone Title/Abstract}) \text{ OR (high-density polyethylene} \text{ Title/Abstract}) \text{ OR (Medpor Title/Abstract}) \text{ OR (polytetrafluoroethylene} \text{ Title/Abstract}) \text{ OR (Gore-Tex} \text{ Title/Abstract}) \text{ AND (((rhinoplasty} \text{ Title/Abstract}) \text{ OR (nasal augmentation} \text{ Title/Abstract}) \text{ OR (revisional rhinoplasty} \text{ Title/Abstract}) \text{ OR (dorsum augmentation} \text{ Title/Abstract}) \text{ OR (nasal dorsum} \text{ Title/Abstract})\) |
| Google Scholar (172,11) \(\text{Concept 1: allintitle: "alloplast" OR "silicone" OR "high-density polyethylene" OR "Medpor" OR "polytetrafluoroethylene" OR "Gore-Tex" "nasal dorsum"  
Concept 2: allintitle: "alloplast" OR "silicone" OR "high-density polyethylene" OR "Medpor" OR "polytetrafluoroethylene" OR "Gore-Tex" "Rhinoplasty" \) |
| Cochrane library (27) \((\text{alloplast}) \text{ OR (silicone) OR (high-density polyethylene}) \text{ OR (Medpor) OR (polytetrafluoroethylene}) \text{ OR (Gore-Tex}) \text{ AND ((rhinoplasty}) \text{ OR (nasal augmentation}) \text{ OR (revisional rhinoplasty}) \text{ OR(dorsum augmentation}) \text{ OR (nasal dorsum})\) |

![Fig. 1 PRISMA Flowchart for study selection](image)
Table 2  Characteristics of the included studies

| Author (year), country of origin | Study type | Mean follow-up (range) | Number of cases (primary/secondary/tertiary) | Mean age/sex | Satisfaction |
|----------------------------------|------------|------------------------|---------------------------------------------|--------------|--------------|
| Yap, E. C. et al. (2011), Philippines | Retrospective | Initial follow-up was on the fifth to seventh postoperative day. Successive follow-up visits occurred 2 weeks, 6 months, and 1 year after surgery | 1054 (1008 primary, 46 secondary) | 34 (15 to 72)/955 female and 99 male | 99.62% |
| Kim, Y. S. et al. (2015), Korea | Retrospective | 9 to 108 months (mean 293 months) | 11 (all secondary) | 37.1 years (8 female 3 male) | 81.81% |
| Scott Shadfar et al. (2015), Pennsylvania | Retrospective | 9(1–47) | 35 (23 primary, 12 secondary) | 36 (17 to 65)/NR | Not mentioned |
| Joo, Y. H. et al. (2016), Republic of Korea | Retrospective | 12(4–115) | 176 (17 revision) | 30.3 (11 to 69)/(96 male, 80 female) | 75% |
| Winkler, A. A. et al. (2012), USA | Retrospective | 12.1(0–74) | 75 | 46 (7 to 86) | Not mentioned |
| Beekhuis, G. J. et al. (1980), USA | Retrospective | NR | 30 | NR | Not mentioned |
| Alvarez-Buylla Blanco, M et al. (2011), Spain | Retrospective | Up to 12 years | 30(NR) | NR | Not mentioned |
| Karnes, J et al. (2000), USA | Retrospective | NR | 93 | NR | Not mentioned |
| Niechajev, I (2012), USA | Retrospective | 6 months to 15 years (median, 7 years) | 52 | 18 to 70 years (median, 29) | 90.56% |
| Han (2012), South Korea | Prospective/cohort | 20.9 months (2–105 months) | 58 | 29.4y(14–62 years)/5 male, 53 female | nm |
| Chen (2010), Taiwan | Retrospective | 254 m (5–71) | 32 | 22 years (16–31) | 90.6% |
| Hong et al. (2010), South Korea | Prospective/cohort | 34 months (12–98 months) | 873 total | 24 years (18–57) | 84.4% patient satisfaction |
| Schwaiger et al. (2019), UK | Retrospective | 342 months (1–106 months) | 51 total case | 24 male, 27 female | nm |
| Jeong et al. (2018), South Korea | Retrospective | 6 months | 227 | 25 years (22–38 years)/21 male, 206 female | 91.2% |
| Turegun. M et al. (2008), Turkey | Prospective | 30 months | 14 | 35.5 (21–50) | 100% |
| Conrad K et al. (2009), Canada | Retrospective | 71 months (1–17 years) | 349 | – | 94.8% |
| Lohuis. P.F.M et al. (2001), Netherlands | Retrospective | 17.9 months (3–72 months) | 66 | 35.9 years (10–66) years | 23 male and 44 female |
| Author (year), country of origin | Study type | Mean follow-up (range) | Number of cases (primary/secondary/tertiary) | Mean age/sex | Satisfaction |
|---------------------------------|------------|------------------------|---------------------------------------------|--------------|--------------|
| Mohammadi Sh et al. (2014) [29], Iran | Prospective/cohort | 3 years In monthly intervals | 38 | 36 (15–58) years, 39 female and 25 male | Not mentioned |
| Waldman S R et al. (1991) [30], USA | Retrospective | – (12–36) months | 17 | 33 years (17–48) years 10 female and 7 male | 94.1% |
| Zelken Jonathan et al. [31] (2017), Taiwan | Retrospective | 6 months (1–36) months | 177 | 34 years (19–72) years 159 female and 18 male | 4 unsatisfied |
| Godin. M et al. [32] (1995), USA | Retrospective | 25 months (6–80) months | 137 | 36 years (14–68) years Sex is not mentioned | 100% All 137 patients |
| Hwan Wang J et al. [33] (2007), Korea | Retrospective | 31 months (12–39) months | 27 | 33 years (16–65) years 15 female and 12 male | 88.8% |
| Zeng Yanjun et al. [34] (2002), China | Prospective | – 3 months–5 years | 98 | 63 (17–49) years 77 female and 21 male | 63% |
| Pham (2011) [35] USA | Retrospective | 36 months | 23 | 20–57 years 1 male 22 female | 1 not satisfied |
| Pham and Hunter [36] (2006) USA | Retrospective | 3 months–5 years | 19 | 18–56 years 19 female 0 male | 1 not satisfied |
| Niechajev [37] 1999 Sweden | Prospective | 1–3 years | 23 dorsal | 30 years (23–47 years) 16 female 11 male | nm |

**Abbreviations:** nm not mentioned
| Study, year of publication (number of cases) | Study design | Rates of complications | Revision |
|-------------------------------------------|-------------|-----------------------|----------|
| Yap. E. C. et al. [11] 2011 (n = 1054)    | e-PTFE      | Extrusion: 5  Infection: 4  Deviation (graft displacement): 11  Overcorrection: 4  Hematoma: 0  Irregularity: 0  Others: 8  | 8 |
| Turegun. M et al. [26] 2008, (n = 14)     | Medpor      | Extrusion: 0  Infection: 0  Deviation (graft displacement): 1  Overcorrection: 0  Hematoma: 0  Irregularity: 0  Others: 3 patients indicated that the difficulty of breathing started right after the surgery and 11 patients stated that it became worse.  | 1 |
| Conrad. K et al [27] 2009, (n = 349)      | Gore-Tex    | Extrusion: 2  Infection: 0  Deviation (graft displacement): 0  Overcorrection: 0  Hematoma: 0  Irregularity: 0  Others: 4 Soft tissue reaction  | 20 |
| Lohuis. P.J.F.M et al. [28] 2001, (n = 66) | Gore-Tex    | Extrusion: 0  Infection: 0  Deviation (graft displacement): 1  Overcorrection: 0  Hematoma: 0  Irregularity: 0  Others: 0  | 1 |
| Mohammadi Sh et al. [29] 2014, (n = 38)   | Medpor      | Extrusion: 0  Infection: 2  Deviation (graft displacement): 0  Overcorrection: 0  Hematoma: 0  Irregularity: 0  Others: 0  | 1 |
| Waldman S R et al [30] 1991, (n = 17)     | Gore-Tex    | Extrusion: 0  Infection: 0  Deviation (graft displacement): 1  Overcorrection: 0  Hematoma: 0  Irregularity: 0  Others: 0  | 1 |
| Zelken Jonathan et al. [38] 2016, (n = 177) | Composite (silicone-PTFE) | Extrusion: 0  Infection: 2  Deviation (graft displacement): 8  Overcorrection: 0  Hematoma: 4  Irregularity: 0  Others: 0  | 12 |
| Godin. M et al [32] 1995, (n = 137)       | Gore-Tex    | Extrusion: 0  Infection: 0  Deviation (graft displacement): 1  Overcorrection: 0  Hematoma: 0  Irregularity: 0  Others: 4  | 4 |
| Hwan Wang J et al. [33] 2007, (n = 27)    | silicone    | Extrusion: 1  Infection: 1  Deviation (graft displacement): 0  Overcorrection: 0  Hematoma: 0  Irregularity: 0  Others: 0  | 3 |
| Zeng Yanjun et al. [34] 2002, (n = 98)    | silicone    | Extrusion: 0  Infection: 38  Deviation (graft displacement): 0  Overcorrection: 15  Hematoma: 0  Irregularity: 0  Others: Drift of prosthesis Severe: 32 Mild: 43 Convexo-concave at nasal root Small: 19 Significant: 7 Small angle of the nose bridge less than 25°: 27  | Nm |


| Study, year of publication (number of cases) | Study design | Rates of complications | Revision |
|---------------------------------------------|--------------|------------------------|----------|
|                                             |              | Extrusion | Infection | Deviation (graft displacement) | Overcorrection | Hematoma | Irregularity | Others |                |
| Hong et al. [23], 2010 (n = 257) | Prospective/cohort | 0 | 2 | 3 | 3 | 0 | 8 | Skin thinning 1/tip problems 9/ minor problems 4/ 2 too low dorsum / 34 |
| Schwager et al. [24] (2019) (n = 20) | Retrospective | 0 | 1 | 2 | 0 | 0 | 0 | 0 | Nm |
| Chen et al. [22], 2010 (n = 32) | Retrospective | 2 | 2 | 0 | 0 | 0 | 0 | 0 | 2 |
| Han et al. [21], 2012 (n = 58) | Prospective/cohort | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 2 |
| Jeong et al. [25] (2018) (n = 227) | Retrospective | 2 | 5 | 4 | 0 | 0 | 7 | 0 | 7 |
| Niechajev, I [20], 2012 (n = 53) | Medpor | 2 | 3 | 0 | 2 | 0 | 0 | 1 building 1 insufficient augmentation Patient dissatisfaction = 5 |
| Colton, J.J et al. [19], 1992 (n = 93) | Mersilene | 0 | 8 | 0 | 0 | 0 | 0 | 0 | 4 |
| Karnes Julie et al. 2000 [17] (n = 30) | Medpor | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| Alvarez-Buylla Blanco, M et al. [17], 2011 (n = 14) | Gore-Tex | 0 | 4 | 0 | 0 | 0 | 0 | 0 | 3 |
| Beekhuis, G. J et al. [16], 1980 (n = 30) | Polyamide | 0 | 3 | 0 | 0 | 0 | 0 | 0 | 3 |
| Winkler, A. A. et al. [15], 2012 (n = 75) | ePTFE | 2 | 4 | 0 | 0 | 0 | 0 | 0 | nm |
### Table 3 (continued)

| Study, year of publication (number of cases) | Study design | Rates of complications | Revision |
|--------------------------------------------|--------------|------------------------|----------|
| Joo, Y. H. et al. [14], 2016 (n = 176) ePTFE | Retrospective | Extrusion 0, Infection 1, Deviation (graft displacement) 2, Overcorrection 0, Hematoma 0, Irregularity 1 | Obvious implant contour = 2, Short nose deformity = 1 |
| Scott Shadfar et al. [13], 2015 (n = 35) ePTFE | Retrospective | Extrusion 1, Infection 1, Deviation (graft displacement) 0, Overcorrection 0, Hematoma 0, Irregularity 1 | 0, 2 |
| Kim, Y. S. et al. [12] 2015 (n = 11) 8 silicone 3 Gore-Tex | Retrospective | Extrusion 0, Infection 1, Deviation (graft displacement) 0, Overcorrection 0, Hematoma 0, Irregularity 0 | 0, 2 |
| Pham and Hunter [36] 2006 (n = 19) Medpor | Retrospective | Extrusion 0, Infection 0, Deviation (graft displacement) 0, Overcorrection 0, Hematoma 0, Irregularity 0 | 0, 1 |
| Pham [35] 2011 (n = 23) Medpor | Retrospective | Extrusion 0, Infection 0, Deviation (graft displacement) 0, Overcorrection 0, Hematoma 0, Irregularity 0 | 0, 1 |
| Niechajev [37] 1999 Medpor (n = 23) | Prospective | Extrusion 1, Infection 1, Deviation (graft displacement) 0, Overcorrection 0, Hematoma 0, Irregularity 1 | 2 manageable complications, 2 |
ePTFE, and high-density polyethylene are an alternative. They are associated with varying incidences of infection and extrusion. Owing to their affordability, lack of any graft harvesting site and being tailorable to a particular deformity, in certain circumstances, alloplastic materials might be used [5]. In 2008, Peled et al. conducted a meta-analysis on rates of infection, extrusion, revision, and removal of different implants used in rhinoplasty surgery and mentioned that alloplastic implants have acceptable complication rates and might be used when facing limitations in using autogenous materials [7].

The aim of these meta-analyses is to explore the incidence of complications after dorsum augmentation surgery using alloplastic materials.

**Materials and methods**

**Protocol and registration**

This meta-analysis was registered at the International Prospective Register of Systematic Reviews (https://www.crd.york.ac.uk/PROSPERO, registration number CRD42020209644). Also, the PRISMA 2020 Guidelines were followed in this systematic review and meta-analysis [8].

**PICO question**

(P) Patient: patients with nasal dorsum deformities undergoing reconstructive or cosmetic rhinoplasty. (I) Intervention: reconstructive or cosmetic rhinoplasty of nasal dorsum augmentation without other nasal deformities. (C) Comparison: polymer-based alloplastic materials such as silicone, high-density polyethylene (Medpor), and polytetrafluoroethylene (Gore-Tex). (O) Outcome: complication rates including visible bulging of the graft, hematoma, graft displacement, irregularity, supra-tip depression, infection, deviation, overcorrection, insufficient augmentation, and major resorption.

**Search strategy**

An electronic survey was conducted using the following databases up to and including September 2020 written in English without any time restriction: PubMed/MEDLINE, Google Scholar and the Cochrane Central Register of...
Controlled Trials (Central). The searching was completed by a manual hand search of the references of all selected full-text articles. The following search terms were screened with its appearance limited to title of the article: (a) “rhinoplasty,” (b) “nasal augmentation,” (c) “revisional rhinoplasty,” (d) “dorsum augmentation,” (e) “nasal dorsum,” (f) “alloplast,” (h) “silicone,” (i) “high-density polyethylene,” (j) “Medpor,” (k) “polytetrafluoroethylene,” and (l) “Gore-Tex” (Table 1).

Study selection
Inclusion criteria were as follows:

1. Randomized clinical trials (RCTs), controlled clinical trials (CCTs), prospective and retrospective cohort studies, and case series with more than 10 participants which provided detailed report on complications (visible bulging of the graft, hematoma at the recipient area, graft displacement, irregularity, supra-tip depression, infection, deviation, overcorrection, insufficient augmentation, major resorption). (Report of at least one complication and revision surgery was mandatory.)

2. No follow-up restrictions
3. Only papers in English are included

Exclusion criteria were as follows:

1. Any cadaver studies or nonhuman studies
2. Studies reporting ratios (risk ratio, odds ratio, hazard ratio) instead of the absolute outcomes were not of our interest.
3. Any article that did not provide any detailed data regarding complication rates
4. Reports of using graft in other parts than nasal dorsum
5. Reports of using liquid alloplastic materials

Data extraction
Based on a predefined paper checklist, the following data was retrieved from the finally included studies by two reviewers (M A.V and R.G) independently and supervised by third author (Sh.R). Any disagreements were resolved by discussion with a third author (Sh.R).

![Fig. 3 Deviation rates reported for synthetic materials in both fixed and random-effects model](image-url)
Data extraction included the following categories:
First author, year of publication, study location, study type, mean age, mean follow-up (range), sex, number of total cases, and cases with complication, incidence of complications after dorsum augmentation with polymer-based alloplastic materials such as silicone, high-density polyethylene (Medpor), and polytetrafluoroethylene (Gore-Tex), rates of complications, revision surgical procedures, and satisfaction rate (percent). The complications assessed were as follows:
visible bulging of the graft, hematoma at the recipient area, graft displacement, irregularity, supra-tip depression, infection, deviation, overcorrection, insufficient augmentation, major resorption.

Risk of bias assessment within the studies
The methodological quality and synthesis of included materials was assessed using a tool for bias assessment in case series by Murad et al. [9]. There were 8 questions in the following domains: selection, ascertainment, causality, and reporting.

Data analysis
Considering the challenges with meta-analysis in observational studies [10], we carefully checked whether included materials in hand were able to answer our clinical question (PICO). The proportion meta-analysis was performed using MedCalc version 18.9.1 (MedCalc Software Ltd., Ostend, Belgium). Both random and fixed model were used based on the heterogeneity. If the heterogeneity was significant, random model was preferred. We conducted the \( \chi^2 \) and \( I^2 \) tests to convey the potential heterogeneity. Potential publication biases were evaluated using funnel plots.

Results
Study selection
Figure 1 shows the PRISMA flow diagram for the study selection process at different stages. 572 papers were obtained through the first search. After duplication removal, 491 papers remained that title and abstract were assessed for eligibility. Reports sought
for retrieval of 77 papers. Of those, 50 papers were excluded with reason (3 reports not retrieved) and finally 27 papers remained which were included in the analysis [11–37].

Study characteristics
The characteristics of included materials are shown in Table 2. Regarding the study type, 27 observational studies were included, 21 retrospective, and 6 prospective case series. A total of 3803 cases were enrolled in this systematic review and meta-analysis. The mean age of patients was 33 (age range 10–72). Although 3 papers did not specify mean and/or age range [16–19]. Four hundred twenty-eight cases were male and 2573 were female although 10 studies did not specify gender [13–32]. The mean follow-up time was 30 months with a range of 3 months to 15 years. Two papers did not report mean follow-up time specifically [16–19]. The included materials were conducted between years 1980 and 2019 in the following countries: South Korea [12–33], the USA [13–37], Spain [17], Taiwan [22–31], the UK [24], Turkey [26], Canada [27], Netherlands [28], China [34], Iran [29], Sweden [37], and Philippines [11].

Complications
The data on each complication are available in Table 3. The meta-analyses were available for the following complications: infection, deviation, irregularity, hematoma, extrusion, and over correction.

Twenty-seven articles with a sample size of 3153 reported on the incidence of infection after dorsum augmentation with synthetic materials. In a random-effects model, the weighted mean percentage was 2.75% (95% CI 1.61 to 4.17%) (Fig. 2). The same articles (3153 cases) also reported on the deviation and irregularity and extrusion rates; the weighted mean percentage were 1.91% (95% CI 0.77 to 3.54%) (Fig. 3), 0.72% (95% CI 0.316 to 1.31%) (Fig. 4), and 0.78% (95% CI 0.43 to 1.24%) (Fig. 5) respectively. The weighted mean of hematoma and
over-correction in a random-effects model were 0.70% (95% CI 0.24 to 1.40%) and 0.49% (95% CI 0.28 to 0.77%) respectively (Figs. 6 and 7).

Other complications
Some rare complications did not meet the criteria for meta-analysis and therefore reported narratively:

One case of opening of the tube the diced cartilage pieces, in a cleft lip patient, pleural tear, and air leak during rib harvesting, of strike skin necrosis (Table 3)

Revision rates
All included material with a total of 451 patients reported on revision surgery rates; the pooled rate was 6.40% (95% CI 3.81 to 9.57%) (Fig. 8). Four papers did not report a specific number of revision surgery and therefore not included in the meta-analysis. The revision rates for the three most commonly used materials (Medpore, Gore-Tex, and silicone) were 6.61% (95% CI 3.98 to 9.85%), 4.91% (95% CI 1.81 to 9.43%), and 7.64% (95% CI 4.93 to 10.88%) respectively (Figs. 9, 10, and 11).

Publication bias
We performed funnel plot for publication bias assessment for each of variables. In the current study, some levels of bias were reported for all complications.

Discussion
One of the greatest challenges in rhinoplastic surgeries is the management of nasal dorsum augmentation. Due to the ease of use, producing ideal aesthetic results and removing needed for graft harvesting sites, alloplastic materials play an important role in typical cosmetic dorsal augmentation [1]. The use of alloplastic materials to address dorsal deficiencies is common among patients avoiding autogenous tissue harvest. Also, patients with several prior nasal operations and significant deformities are the best candidates for alloplastic materials as they mostly have depleted potential autogenous harvesting site [7]. Although there are concerns over several complications associated with synthetic materials including infection, deviation, extrusion, etc. [1]. In this systematic review and meta-analysis, we determined the

![Fig. 6 Hematoma rates reported for synthetic materials in both fixed and random-effects model](image-url)
complications reported for different alloplastic materials. Twenty-seven articles reported on complications and outcomes of dorsal augmentation rhinoplasty with synthetic materials. The widely used alloplasts were expanded polytetrafluoroethylene (ePTFE), high-density polyethylene, and silicone. The total rates for complications, infection, deviation, irregularity, hematoma, extrusion, and over correction were 2.75%, 1.91%, 0.72%, 0.70%, 0.78%, and 0.49%, respectively. The revision rate, based on random effects model, was 6.40% with 95% CI (3.84 to 9.57).

We reported a subsequent revision of 0–21% in our included studies. The pooled rate for the need for revision surgery was 6.40%. The revision rates for the three most commonly used materials Med-pore, ePTFE, and silicone were 6.61%, 7.06%, and 7.64%, respectively. The decision for implant removal is quite controversial; although, surgical removal of infected implants followed by an immediate or delayed reconstruction has higher chances of resolution [39].

The highest revision rates were related to silicone (7.64%). A similar study reported 6.5% revision rate for silicone implants [7]. Being the most commonly used alloplastic material in Asian countries, silicone is a smooth, cost benefit, and easy-carved implant which can be easily removed in case of failure. The lack of pores leads to fibrous capsule formation around the implant within the body.

Infections and displacement are the main causes of revision surgery in silicones and therefore in order to reduce such problems aggressive modification of the natural barriers and anatomical structure should be strictly avoided [40]. If shaped appropriately according to the nasal phenotype, the extrusion rate would reduce [41]. To manage and reduce complications, this method supports alloplastic materials better for patients with thicker skin than for patients with thinner skin.

The high-density polyethylene (Medpore), with pore size range from 160 to 368 μm, and more than half of these pores are larger than 150 μm in diameter and have
excellent biocompatibility. In candidates of augmentation rhinoplasty with severe over resections or severe deformities, these implants have been a useful option. Our findings for revision surgery of high-density polyethylene have been higher than previously reported rates [7].

The revision rate for polytetrafluoroethylene/expanded polytetrafluoroethylene (Gore-Tex) was 4.91%. This hydrophobic polymer with pores of up to 30 μm allows for bacterial adherence and levels of issue integration that provides implant stability with ease of removal if needed. Our results are in line with previous studies mentioning low incidence of revision rates compared with other synthetic materials. A similar previous meta-analysis (in 2008) reported the removal of 3.1% for both ePTFE and high-density polyethylene [7].

Nevertheless, our results suggested a relatively high total rate for revision rate (6.40%) compared with autogenous grafts (3.03%) [42]. This might be attributed to the fact that infection in synthetic materials, unlike autogenous grafts, conservative treatments are inapplicable and mostly require revisional surgery [42, 43].

The use of autograft materials in nasal dorsum augmentation is a safer treatment with fewer complications compared to the alloplastic method. Complications of using autogenous grafts materials such as diced cartilage include graft resorption, insufficient augmentation, deviation (graft displacement), infection, irregularity, supra-tip depression, over-correction, hematoma at the recipient site, and the visible bulging of the graft. According to the findings of the article, infections caused by the use of alloplastic usually require revision surgery, while most infections that occur in the autograft method can be controlled by intravenous antibiotics [42].

In fact, it can be said that the complications of the autograft method are manageable and controllable complications.

Also, the use of alloplastic materials is a risky method in comparison with autograft materials taken from the patient himself, because the use of alloplastic materials acts as a foreign body in the body and its high-risk side effects can lead to nasal deformity and aesthetic complications. Undesirability in the systematic study [44]
Fig. 9 Revision rates reported for Medpore in both fixed and random-effects model.

Fig. 10 Revision rates reported for Gore-Tex in both fixed and random-effects model.
was reported in autograft materials, which are usually removed from the abdomen or thighs, has fewer reported complications after surgery, and in most cases, complications such as numbness gradually decrease after surgery and are completely eliminated by 3 months after surgery.

In addition to the side effects of using alloplastic materials, some side effects may be preventable, such as bending along the natural convexity, bone resorption, and foreign body reactions such as fibrous capsule formation and tissue ingrowth.

Due to the fact that the complications mentioned throughout the article are not only common but also lead to major problems both during surgery and after surgery. By examining the problems and complications of this method, treatment-related techniques will be developed in the future. With the passage of time and the development of new surgical methods and materials, it shows that current methods are always associated with complications, and at no time are failures and complications announced at the same time as successes.

Current articles widely support autografts instead of using alloplastic in rhinoplasty. Surgeons describe alloplastic implants as dangerous, unpredictable, and hard to use. Therefore, the reported complications are less than 5%. Complications that require revision surgery and cause the material to be removed are 3.7%, which is a significant amount compared to the use of autografts [45], which is 1% [96]. Therefore, the use of alloplastic materials seems to be mentioned with the desire of the patient and the surgeon and acceptance of the possibility of complications (Tables 4 and 5).

Limitations and strengths
Reports of complications often come from other investigators, citing their own experience with implants inserted by other surgeons. The major limitation with current systematic review and meta-analysis was the descriptive nature of much of the current literature and lacking comparator groups. Also, a proper tool for quality assessment in case series was lacking and we had to make adaptations in domains. Some levels of bias might be caused by excluding non-English materials.

The time frame for follow-ups in this study assumed rationale for complications appearance. Having a clear understanding of complications of each material and the ways to prevent and treat them is possible by accurate disclosure of shortcoming in the literature. In the future, the development of alloplasts that approximate the ideal implant with low complication rates is warranted. The technology of prefabrication of precise three-dimensional bioactive and biocompatible implants might reduce the incidence of complications and lower the chance of failure.

Conclusion
To recapitulate, this meta-analysis suggested an acceptable rate of complications and revision surgery with synthetic materials. Synthetic materials might be a proper
Table 4  Excluded studies with reason

| Title (reference)                                                                                                                                                                                                 | Reason on exclusion |
|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------|
| 1. Medpor in maxillofacial deformities: report of three cases [46]                                                                                                                                                 | 3 cases             |
| 2. A case report of ophthalmic artery emboli secondary to calcium hydroxylapatite filler injection for nose augmentation—long-term outcome [47]                                                                | Not solid polymers  |
| 3. Case reports of adipose-derived stem cell therapy for nasal skin necrosis after filler injection [48]                                                                                                          | Not solid polymers  |
| 4. Complete septal extension grafts using porous high-density polyethylene sheets for the westernization of the Asian nose. [21]                                                                             | Not for dorsum      |
| 5. Retinal branch artery embolization following hyaluronic acid injection: a case report [49]                                                                                                                     | Not solid polymers  |
| 6. Two cases of adverse reactions of hyaluronic acid-based filler injections [50]                                                                                                                                  | Not solid polymers  |
| 7. A newly designed minigraft to achieve angulatory and projection of the nasal tip: the asymmetrical bipyramidal graft [51]                                                                                         | Not dorsum separately |
| 8. Foreign body reaction to Radiesse: 2 cases [52]                                                                                                                                                                 | 2 cases             |
| 9. Plastic surgery for women [53]                                                                                                                                                                               | Not related         |
| 10. Midline volume filler injection for facial rejuvenation and contouring in Asians [54]                                                                                                                      | Not solid polymers  |
| 11. Non-surgical rhinoplasty with hyaluronic acid fillers: predictable results using software for the evaluation of nasal angles [55]                                                                       | Not solid polymers  |
| 12. Nasal filling in plastic surgery practice: primary nasal filling, nasal filling for post-rhinoplasty defects, rhinoplasty after hyaluronidase injection in dissatisfied nasal filling patients [56] | Not solid polymers  |
| 13. Calcium hydroxylapatite gel (Radiesse) injection for the correction of postrhinoplasty contour deficiencies and asymmetries [57]                                                                        | Not dorsum separately |
| 14. Augmentation rhinoplasty: observations on 1200 cases [58]                                                                                                                                                   | Not solid polymers  |
| 15. Secondary rhinoplasty of the Asian nose: correction of the contracted nose [59]                                                                                                                              | Not solid polymers  |
| 16. Revision rhinoplasty in ethnic patients: polybeak deformity and persistent bulbous tip [60]                                                                                                               | Not dorsum separately |
| 17. Correction of the supratip deformity of the nose [61]                                                                                                                                                         | Not dorsum separately |
| 18. Assessment of nostril symmetry after primary cleft rhinoplasty in patients with complete unilateral cleft tip and palate [62]                                                                                 | Complications were not assessed |
| 19. Operative techniques in Asian rhinoplasty [63]                                                                                                                                                               | Operative techniques |
| 20. E-M shaped septal encircling with Medpor reconstruction on crooked noses: personal technique and postoperative results [64]                                                                               | Septal encircling reconstruction, not related |
| 21. Late complications of nasal augmentation using silicone implants [65]                                                                                                                                         | Complications were not mentioned |
| 22. Periorbital necrotizing fasciitis and orbital apex syndrome as a delayed but emergent complication of silicone nasal augmentation [66]                                                           | Case report         |
| 23. Management of wide nasofrontal angle with GORE-TEX implants [67]                                                                                                                                            | Not dorsum          |
| 24. Silicone rubber implants in nasal reconstructive surgery [68]                                                                                                                                               | Not dorsum separately |
| 25. Availability and safety of osteotomy in esthetic rhinoplasty of east Asian patients [25]                                                                                                                     | Not dorsum separately |
| 26. Evaluation and proportion in nasal filling with hyaluronic acid [69]                                                                                                                                          | Not found yet. Searching (not retrieved) |
| 27. Prevention and management of iatrogenic blindness associated with aesthetic filler injections [70]                                                                                                                  | Not related         |
| 28. Efficacy and safety of a hyaluronic acid filler to correct aesthetically detracting or deficient features of the Asian nose: a prospective, open-label, long-term study [71] | Not solid polymers  |
| 29. Application of a porous polyethylene spreader graft for nasal lengthening in Asian patients [72]                                                                                                          | Not dorsum separately |
| 30. Use of fillers in rhinoplasty [73]                                                                                                                                                                          | Not solid polymers  |
| 31. Simple implant augmentation rhinoplasty [74]                                                                                                                                                              | No number           |
| 32. Soft and firm alloplastic implants in rhinoplasty: why, when and how to use them: a review of 311 cases [75]                                                                                               | Not dorsum separately |
| 33. The use of expanded polytetrafluoroethylene in short nose elongation: fourteen years of clinical experience [76]                                                                                           | L shaped            |
| 34. A novel method to enhance dynamic rhinoplasty outcomes: double “V” carving for alloplastic grafts [77]                                                                                                     | L-shaped            |
| 35. The nonsurgical rhinoplasty: a retrospective review of 5000 treatments [78]                                                                                                                                  | Not dorsum separately |
| 36. The use of Medpor implants for midface contouring in cleft patients [24]                                                                                                                                     | Not dorsum separately |
| 37. Long-term results of high-density porous polyethylene implants in facial skeletal augmentation: an Indian perspective [79]                                                                                 | Not dorsum separately |
| 38. Are polytetrafluoroethylene (Gore-Tex) implants an alternative material for nasal dorsal augmentation in Asians? [23]                                                                                         | Not dorsum separately |
### Table 4  (continued)

| Title (reference) | Reason on exclusion |
|-------------------|----------------------|
| 39. Nasal dorsum reconstruction with alloplastic material [80] | Complications were not mentioned |
| 40. Injection rhinoplasty with hyaluronic acid and calcium hydroxyapatite: a retrospective survey investigating outcome and complication rates [81] | 3 cases |
| 41. Use of porous high-density polyethylene in revision rhinoplasty and in the platyrrhine nose [82] | Not dorsum separately |
| 42. Soft tissue fillers in the nose [83] | Not solid polymers |
| 43. Problems associated with alloplastic materials in rhinoplasty [84] | Complications not mentioned |
| 44. [Nasal dorsal augmentation] [85] | Not in English (not retrieved) |
| 45. Rhinofilling with hyaluronic acid thought as a cartilage graft [86] | Not solid polymers |
| 46. A simple technique for the correction of maxillonasal dysplasia using customized expanded polytetrafluoroethylene (ePTFE) implants [87] | “L”-shaped ePTFE |
| 47. Revision rhinoplasty of Asian noses: analysis and treatment [88] | Complications not mentioned |
| 48. [Pyodermatitis of the nasal pyramid disclosing a complication of rhinoplasty with silicone implant] [89] | Not in English (not retrieved) |
| 49. Use of porous high-density polyethylene in revision rhinoplasty and in the platyrrhine nose (Romo III et al.) [82] | 68 patients had dorsum tip implants/complications were not categorized |
| 50. Nonsurgical rhinoplasty with the novel hyaluronic acid filler VYC-25L: results using a nasal grid approach (Bertossi et al.) [90] | Not solid polymers |

### Table 5  Risk of bias assessment

| Yap, E. C. et al. (2011), [11] Philippines | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | Total |
|-------------------------------------------|---|---|---|---|---|---|---|---|------|
| Kim, Y. S. et al. (2015) [12], Korea    | y | y | y | NA | NA | NA | Y | y | 5/8  |
| Scott Shadfar et al. (2015) [13], Pennsylvania | y | y | y | NA | NA | NA | Y | y | 5/8  |
| Joo, Y. H. et al. (2016) [14], Republic of Korea. | y | y | y | NA | NA | NA | Y | y | 5/8  |
| Winkler, A. A. et al. (2012) [15] USA. | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Beekhuis, G. J. et al. (1980) [16], USA | y | y | y | NA | NA | NA | N | y | 4/8  |
| Alvarez-Buya Blanco, M et al. (2011) [17], Spain | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Karnes, J. et al. (2000) [18], USA | y | y | y | NA | NA | NA | Y | y | 5/8  |
| Colton, J. J. et al. (1992) [19], USA | y | y | Y | NA | NA | NA | N | y | 4/8  |
| Niechajev, I (2012) [20], USA | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Han (2012) [21] South Korea | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Chen (2010) [22] Taiwan | y | y | y | NA | NA | NA | Y | y | 5/8  |
| Hong et al. (2010) [23] South Korea | y | y | y | NA | NA | NA | Y | y | 5/8  |
| Schaeger et al. (2019) [24] UK | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Jeong et al. (2018) [25] South Korea | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Turegun, M et al. (2008) [26] Turkey | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Conrad. K et al. (2009) [27] Canada | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Lohuis. P.J.F.M et al. (2001) [28] Netherland | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Chen Liang et al. (2014) [91] China | y | y | y | NA | NA | NA | Y | y | 5/8  |
| Mohammadi Sh et al. (2014) [29] Iran | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Waldman S R et al. (1991) [30] USA | y | y | y | NA | NA | NA | Y | y | 5/8  |
| Zeiken Jonathan et al. et al. [31] (2017) Taiwan | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Godin. M et al. [32] (1995) USA | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Hwan Wang J et al. [33] (2007) – Korea | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Zeng Yanjun et al. et al. [34] (2002) China | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Pham (2011) [35] USA | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Pham and Hunter [36] (2006) USA | y | y | Y | NA | NA | NA | Y | y | 5/8  |
| Niechajev [37] 1999 Sweden | y | y | y | NA | NA | NA | Y | y | 5/8  |
option when the use of autogenous grafts is not applicable. Judicious case selection and prompt management of complications are crucial when alloplastic materials are used. Some practical clinical recommendations may be helpful in future research and clinical procedures. These recommendations are just based on experts’ experience.

**Abbreviations**
ePTFE: Polytetrafluoroethylene; PRISMA: Preferred Reporting Items for Systematic Reviews; RCT: Randomized clinical trial; CCT: Controlled clinical trial.

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**Authors’ contributions**
O.K contributed to the design, drafting of the article, data analysis/interpretation, statistics, data collection, and critical revision of the article. Sh.R contributed to the design, data interpretation, drafting of the article, critical revision of the article, approval of article: critical revision of the article and approval of the article. G.Y contributed to the data analysis/interpretation, statistics, and data collection. H.R.F contributed to the critical revision of the article and approval of article. M.P contributed to the critical revision of the article and approval of the article. All authors are the major contributors in writing the manuscript. All authors read and approved the final manuscript.

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The authors declare that they have no competing interests.

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