Research Article

Clinical Efficacy and Safety of Different Dental Prosthetic Membranes in Guided Bone Regeneration during Dental Implants: A Meta-Analysis

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Received 24 May 2022; Revised 23 June 2022; Accepted 4 July 2022; Published 31 July 2022

Academic Editor: Dong Chen

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Objective. To evaluate clinical efficacy and safety of absorbable and non-absorbable dental restorative membranes in guided bone regeneration (GBR). Articles concerning absorbable and non-absorbable prosthetic membrane-related studies of GBR were screened from multiple databases. In the end, 526 postoperative patients who met eligibility criteria were screened for the study from eight trials. The results showed that the repair success rate of the experimental group (absorbable dental restorative membrane) was higher than that of the control group (non-absorbable dental restorative membrane) (RR = 1.18, 95% CI [1.11, 1.26], and the total physical therapy effect was \(P < 0.0001, I^2 = 0\)), and the height of bone graft in the experimental group was higher than that in the control group (MD = 0.67, 95% CI [0.11, 1.23]). The thickness of bone graft in the experimental group was higher than that in the control group (MD = 0.43, 95% CI [0.30, 0.56], \(P < 0.00001, I^2 = 61\)), and the adverse events in the experimental group were less than those in the control group (RR = 0.31, 95% CI [0.18, 0.51], \(P < 0.00001, I^2 = 13\%\)). Absorbable prosthetic membrane is superior to non-absorbable prosthetic membrane in clinical efficacy and safety.

1. Introduction

Guided Bone Regeneration (GBR) originated from the field of periodontology in guided tissue regeneration technology. It is a biofilm made of biomaterials, which erect a biological barrier between bone defects and gingival soft tissue [1–3]. GBR prevents epithelial cells and fibroblasts in soft tissue and soft tissue from growing into the bone defect area. This process ensures that the osteogenesis process is completed on the premise of no interference of fibroblasts. Finally, GBR can realize complete bone repair of the defect area, which needs oral repair membrane [4].

Oral repair membrane is a biocompatible material. The repair membrane is placed between oral soft tissue and bone defect by surgery to establish a biological barrier to create a relatively closed bone regeneration environment [5, 6]. Oral repair film can be divided into absorbable film and non-absorbable film according to whether the material can be degraded. In the past, patients used titanium membrane (non-absorbable membrane) as a protective barrier membrane because titanium membrane could not be fully absorbed. This process limited the supply of blood plasma and then hindered the blood supply in the bone graft area, which had a significant impact on the recovery of patients to a certain extent [7–9]. However, it has good plasticity and can bend, trim the contour, adapt to various bone defect forms, better stabilize the wound, and guide bone regeneration [10].

Although the absorbable membrane risks rapid degradation, it makes the new bone tissue adhere to the biofilm. Absorbable membrane promotes the early tissue integration and the production of transmembrane blood vessels, avoids the inward growth of connective tissue, increases the stability of gingival tissue, and reduces gingival atrophy [11]. Also, it can reduce patient
complications without the need for second-stage surgical removal of the membrane.

Although there are several research studies about comparison between absorbable and non-absorbable dental restorative membrane in guided bone regeneration, there is little comprehensive analysis for the topic. Therefore, we conducted this research to overall analyze the difference in absorbable and non-absorbable dental restorative membrane in guided bone regeneration.

In this paper, we have evaluated clinical efficacy and safety of absorbable and non-absorbable dental restorative membranes in guided bone regeneration (GBR). For this purpose, both absorbable and non-absorbable prosthetic films for GBR were selected from multiple databases (PubMed, Web of Science, Cochrane Library, and China National Knowledge Infrastructure), whereas Review Manager 5.2 was used for meta-analysis, sensitivity analysis, and bias analysis. After the screening process, 526 postoperative patients were extracted from 8 trials which are those patients who finally met the qualification criteria to conduct this meta-analysis.

2. Proposed Method or Strategy

To ensure the scientificity, we followed PRISMA statement and the methods of Cao et al. [12].

2.1. Literature Search Strategy. We have searched the randomized controlled trials published by PubMed, ScienceNet, Cochrane Library, and China National knowledge Infrastructure from January 1, 2000, to September 1, 2021, using the following search terms:

(1) Absorbable dental repair membrane.
(2) Bone regeneration.
(3) Clinical effect. The search strategy involves medical subject headings (mesh) and text words combined by the Boolean operator "AND."

We will conduct a comprehensive search in multiple databases without restrictions on language or publication status. In order to maximize the specificity and sensitivity of the search, the author should also refer to the list of retrieved references to find other relevant studies not found through the search strategy.

A comprehensive review of potentially relevant articles was conducted to ensure that they met all inclusion criteria, as follows:

(1) Studies comparing patients receiving absorbable and non-absorbable dental repair membranes.
(2) Studies comparing patients receiving absorbable and non-absorbable dental repair membranes.
(3) GBR patients.
(4) Between absorbable and non-absorbable dental restorative membranes, indexes for evaluating curative effect or other relevant indexes are included.
(5) The full text is available for reference.

Studies were excluded according to the following predetermined exclusion criteria:

(1) Studies on other subjects.
(2) Comparison of other interventions.
(3) Lack of research on available data.
(4) Comments, abstracts, and reproduction of publications.

2.2. Data Extraction and Quality Assessment. Two pairs of reviewers independently screened the titles, abstracts, and full-text articles of potentially qualified studies and resolved their differences through discussion. The following data parameters were extracted: name of main author, study country, patient population in the study, number of participants in each group, patient age, patient gender, characteristics of drug intervention during follow-up in each group, and outcome measurement in each group. The Cochrane bias risk tool in Review Manager 5.2 was used to evaluate the effectiveness of qualified randomized controlled trials. Egger's test and funnel plot program were used to assess the risk of bias in the study.

2.3. Statistical Analysis. Review Manager (version 5.2, Cochrane Collaboration, 2011) is used to evaluate the impact of the results in the selected report. In order to measure the consistency of effect size (or and MD), DerSimonian and Laird random effect models were used for paired meta-analysis, and the combined estimates and 95% CI between two groups were calculated. 0% to 40% of heterogeneity is considered "may not be important," 30% to 60% is considered "moderate heterogeneity," 50% to 90% is considered "substantial heterogeneity," and 75% to 100% is considered "considerable heterogeneity."

If $P < 0.05$ or $I^2 > 50\%$, the random effect model was used for analysis; if $P \geq 0.05$ and $I^2 \leq 50\%$, the fixed effect model was used for analysis. When heterogeneity exists, the random effect model is used, while the fixed effect model is applied. Publication bias was examined by visual examination of the funnel plot and using Egger's test. Sensitivity analysis was performed by deleting one study at a time to observe the impact of individual results on the overall analysis.

3. Results

3.1. Search Process. The initial search yielded 966 articles from four databases, including PubMed, Embase, Web of Science, and CNKI. After the first screening, 880 records were retained. By screening titles and abstracts, additional 823 records were excluded because they were review articles, letters, case reports, comments, or editorials. Then, 57 articles were remained. Eight articles were further excluded for various reasons, including different research designs or insufficient available data.

Finally, 8 studies [13–20] met the inclusion criteria and were included in this meta-analysis, with a total of 526 patients. The process followed PRISMA guidelines, including the reasons for excluding the study, as shown in Figure 1.
3.2. Characteristics of Included Studies. Table 1 lists the main characteristics of the eight tests. These studies included 526 patients (263 patients in the experimental group and 263 patients in the control group). All 8 articles were published from 2016 to 2020. The sample size is between 26 and 100.

3.3. Results of Quality Assessment. The Cochrane bias risk assessment tool was used to assess the risk of inclusion in the study. Of these 8 articles, only 1 study found a high risk of selection bias, performance bias, detection bias, abrasion bias, reporting bias, and other bias (Figures 2 and 3).

Given the deviation summary, only 1 clue has different deviation. Visual examination of the funnel chart of studies reporting efficiency showed some asymmetry, and the Egger test showed little evidence of publication bias.

3.4. Results of Heterogeneity Test

3.4.1. Heterogeneity Analysis of Successful Repair between Experiment and Control Groups. Meta-analysis of successful repair. The overall results showed that the repair success rate of the experimental group was higher than that of the control
3.4.2. Heterogeneity Analysis of Height of Bone Graft between Experiment and Control Groups. Similarly, the first micturition (min) between the experimental group and the control group was meta-analyzed. The overall result showed that the height of bone transplantation in the experimental group was higher than that in the control group (MD = 0.67, 95% confidence interval [0.11, 0.12], P < 0.0001, I² = 99%, using random effect model) (Figure 5).

3.4.3. Heterogeneity Analysis of Bone Graft Thickness between Experiment and Control Groups. For residual urine, it was reported in 7 studies. The overall results showed that the thickness of bone graft in the experimental group was higher than that in the control group (MD = 0.43, 95% confidence interval [0.30, 0.56], P < 0.00001, I² = 61%, using random effect model) (Figure 6).

3.4.4. Heterogeneity Analysis of Adverse Events between Experiment and Control Groups. To better assess the safety of different therapies, we collected data on adverse events. The overall results showed that the adverse events in the experimental group were less than those in the control group (RR = 0.31, 95% confidence interval [0.18, 0.51], P < 0.00001, I² = 13%, using the fixed effect model) (Figure 7).

3.5. Results of Sensitivity Analysis and Publication Bias. To assess the sensitivity of the articles, we deleted a study to observe the effect of individual outcomes on the overall efficacy of urinary retention. In Figure 4, the result shows I² = 0% high heterogeneity. When Wang’s article [18] was deleted, the results change the most, indicating the robustness of the included study (Figure 8).

We used funnel plots to assess the efficiency of urinary retention. Visual results showed symmetrical shape. The P value of Egger test was 0.218, indicating that there was no publication bias in this study (Figure 9).

### Table 1: Characteristics of eligible studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Year</th>
<th>Country</th>
<th>Groups</th>
<th>Sex (male/female)</th>
<th>Age (years)</th>
<th>n</th>
<th>Years of onset</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basler</td>
<td>2018</td>
<td>Switzerland</td>
<td>Resorbable membrane</td>
<td>11/12</td>
<td>56.6 ± 17.4</td>
<td>12</td>
<td>January 2015 to January 2018</td>
</tr>
<tr>
<td>Cui</td>
<td>2019</td>
<td>China</td>
<td>Resorbable membrane</td>
<td>44/36</td>
<td>47.4 ± 4.25</td>
<td>40</td>
<td>February 2016 to February 2018</td>
</tr>
<tr>
<td>Huang</td>
<td>2016</td>
<td>China</td>
<td>Resorbable membrane</td>
<td>63/37</td>
<td>50.8 ± 1.9</td>
<td>50</td>
<td>August 2013 to February 2015</td>
</tr>
<tr>
<td>Naenni</td>
<td>2016</td>
<td>Switzerland</td>
<td>Resorbable membrane</td>
<td>13/14</td>
<td>51.85 ± 29.7</td>
<td>13</td>
<td>March 2010 and January 2013</td>
</tr>
<tr>
<td>Wang</td>
<td>2018</td>
<td>China</td>
<td>Resorbable membrane</td>
<td>39/37</td>
<td>40.25 ± 18.75</td>
<td>38</td>
<td>July 2017 to July 2018</td>
</tr>
<tr>
<td>Wang</td>
<td>2020</td>
<td>China</td>
<td>Resorbable membrane</td>
<td>49/45</td>
<td>42.4 ± 5.55</td>
<td>47</td>
<td>December 2017 to December 2018</td>
</tr>
<tr>
<td>Yang</td>
<td>2016</td>
<td>China</td>
<td>Resorbable membrane</td>
<td>41/25</td>
<td>45.07 ± 6.5</td>
<td>33</td>
<td>January 2014 to January 2015</td>
</tr>
</tbody>
</table>

Figure 2: Graph of the risk of bias: green = low risk; yellow with question mark = unclear; and red = high risk.
4. Discussion

From our results, we can find that absorbable dental restorative membrane had higher successful repair than non-absorbable dental restorative membrane in guided bone regeneration. In addition, height of bone graft and bone graft thickness were both higher in absorbable dental restorative membrane than non-absorbable dental restorative membrane. In the comparison of safety, absorbable dental restorative membrane was worse than non-absorbable dental restorative membrane. These results showed that absorbable dental restorative membrane was better than non-

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**Figure 3:** Risk of bias for each study, using three colors: green = low risk; yellow with question mark = unclear; and red = high risk.

**Figure 4:** Forest plots for the effects for successful repair in experiment versus control groups.
absorbable dental restorative membrane in clinical effects and safety. These were consistent with Zhang’s study [21] that bone regeneration guided by absorbable biofilm in patients with GBR improves the success rate of dental implantation and has high safety.

With the development of dental implants, implant denture has become one of the conventional treatment methods for repairing dentition defects or deletions, and the methods are constantly simplified [22, 23]. The safety and reliability are gradually improved. Guided bone regeneration (GBR) is often widely used to treat periodontal diseases and repair maxillary sinus defects and bone defects. Oral repair membrane has the characteristics of high efficiency, short time consumption, thick osteogenesis, and high
osteogenesis [24]. The primary function of the oral repair membrane is to prevent epithelial cells and connective tissue cells from entering the regeneration area and create and maintain a space for the unrestrained growth of pluripotent stem cells and osteoblasts [25]. It is widely used in stomatology, such as periodontal mucosa, oral implant, and alveolar surgery.

Non-absorbable membrane (titanium membrane) was a commonly used oral repair material in the past. It has the characteristics of stable space, good resistance strength, and hard texture [26]. It plays a specific role in promoting the growth of bone grafts. However, it also has some adverse effects, such as preventing the excellent absorption of blood by bone graft, prolonging patients’ recovery time, and requiring secondary surgery [27]. The incidence of postoperative complications is high, and the osteogenic effect is poor.

The absorbable membrane has collagen composition similar to periodontal connective tissue, including weak immunogenicity and cytotoxicity [28]. The absorbable membrane can promote the chemotaxis of periodontal ligament (PDL) cells and gingival fibroblasts. In addition, it can encourage hemostasis, is easy to operate, and degrades physiologically. Calcification and ossification can occur when approaching bone [29].

5. Conclusion

In this paper, we have evaluated clinical efficacy and safety of absorbable and non-absorbable dental restorative membranes in guided bone regeneration (GBR). For this purpose, both absorbable and non-absorbable prosthetic films for GBR were selected from multiple databases (PubMed, Web of Science, Cochrane Library, and China National Knowledge Infrastructure), whereas Review Manager 5.2 was used for meta-analysis, sensitivity analysis, and bias analysis. After the screening process, 526 postoperative patients were extracted from 8 trials which are those patients who finally met the qualification criteria. The present study showed that absorbable dental restorative membrane was better than non-absorbable dental membrane both in clinical effects and safety. However, our findings should be carefully considered with caution due to small sample size. Studies in various areas with large study population are essential to further confirm our findings in the future. There are some limitations in this study. Firstly, more indicators should be included, and this could be conducted in the future. Secondly, more research studies from various areas could be analyzed in the next research.

In future, we are keen to extend the proposed study to other domains and diseases preferably in smart healthcare sector.

Data Availability

The datasets used and analyzed during the current study are available from the corresponding author upon reasonable request.

Disclosure

Yan Guo and Linghan Su are co-first authors.

Conflicts of Interest

The authors declare that they have no conflicts of interest.
Authors’ Contributions

Yan Guo and Linghan Su contributed equally to this study. The conception of the paper was completed by Caidi Chen and Yan Liu. The data processing was completed by Caidi Chen, Yan Liu, and Jianxue Li. All authors participated in the review of the paper.

Acknowledgments

This study was supported by the general project of Natural Science Foundation of Gansu Province (20JR10RA006).

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