Porcine Acellular Dermal Matrix: An Alternative to Connective Tissue Graft—A Narrative Review

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Received 23 May 2021; Revised 4 July 2021; Accepted 14 August 2021; Published 6 September 2021

Academic Editor: Paolo Francesco Manicone

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Porcine acellular dermal matrix has recently been introduced in dentistry as an alternative to the gold standard connective tissue graft especially for the use in gingival recession treatments and soft tissue augmentation in implant surgery. Connective tissue grafts are inconvenient and require a second surgical site leading to greater morbidity, longer surgical procedures, and a more painful postoperative phase for the patient. Other options such as allografts have ethical concerns and are less available in Europe. Thus, dental professionals have sought other techniques and materials. Porcine acellular dermal matrix results in periodontal recession treatment with a gain in recession coverage as well as increased keratinized tissue and soft tissue augmentation. This leads to more keratinized mucosa and greater tissue thickness. Many studies have been published using collagen matrices, but a few strictly use porcine acellular dermal matrix, which have been studied in prospective randomized clinical trials with a large number of patients and longer follow-up periods (more than 5 years). Nevertheless, more data are needed to confirm that the porcine acellular dermal matrix is a suitable alternative although its favourable results to date suggest a positive future.

1. Introduction

The porcine acellular dermal matrix (PADM) is a collagen matrix known commercially as mucoderm (Botiss Dental, Berlin, Germany). It has recently been introduced in dentistry as a substitute for the gold standard connective tissue graft (CTG) in periodontal and implant plastic surgery [1–4]. CTGs have limited availability especially when treating multiple gingival recessions. The need for a second surgical site can lead to postoperative pain and increased patient discomfort. There is also a greater risk of bleeding as well as a greater risk of damage to the branches of the palatine artery. Other disadvantages include a longer surgical procedure, greater risk of mucosal necrosis, and the need for hypnosis or anaesthesia especially when extracted from the palate. This has prompted periodontists to employ alternative materials [5–9].

Alternatives include allografts (AlloDerm) which is an acellular dermal matrix originally introduced to treat burn patients in the year 1992 and has since been used in medical and dental reconstructive surgery as a substitute for CTG with no risk of rejection or disease transmission [10, 11]. It consists of an allogenic freeze-dried connective tissue matrix, which has its epidermal layer and cellular components removed keeping its native three-dimensional structure composed of collagen and key extracellular matrix components including fibronectin, proteoglycans, and vascular channels which support cell migration and capillary proliferation. Such materials have raised ethical concerns due to their human origin, and most European countries including Spain have imposed heavy restrictions increasing the popularity of alternatives such as PADM [12–14].

PADM has greater availability than ADM; it can be harvested in bigger quantities. PADM consists of three-dimensional sterilized pure collagen types I and III and elastin, which is a stable tissue matrix derived from porcine dermis without being posteriorly cross-linked artificially or put under any other chemical treatment [15]. PADM also passes through a disinfecting process to eliminate all antigenic and tissue rejection constituents such as
noncollagenous proteins and cells as well as bacteria and viruses from the dermis [16].

In vitro and in vivo investigations proved that PADM can increase growth and proliferation of human gingival fibroblasts, osteoblasts, and endothelial cells revealing a capacity for significant revascularization of the collagen structure during early healing [16–18]. Angiogenesis and the development of blood vessels are fundamental for nutrition, oxygen supply, immune cells, mesenchymal stem cells, and growth factors early in the healing period [19]. Lin et al. showed that PADM facilitates migration, adhesion, and proliferation of periodontal ligament cells and human oral fibroblasts [20].

2. Comparison with Connective Tissue Graft in Recession Treatment

One of the main reasons that collagen matrices have been introduced in dentistry is to avoid using connective tissue grafts for the motives mentioned earlier in this article. Many articles have been published comparing the gold standard with collagen matrices such as PADM for gingival recession treatment (see Table 1). This can increase the width of the attached gingiva on teeth and soft tissue augmentation during implant therapy.

Schmitt et al. published a 10-month preclinical study comparing CTG and PADM in beagle dogs and concluded that PADM was not statistically inferior for tooth volume augmentation of keratinized gingiva than CTG [3].

Cieslìk-Wegemund et al. published a six-month randomized clinical trial comparing PADM with CTG using the tunnel technique to treat Miller Class I and II periodontal recessions stating an adequate recession coverage in patients treated with collagen matrix. There was a gain in keratinized tissue similar to CTG although there were fewer complete recession coverages [2]. Favourable results were also observed in a 12-month follow-up by Cosgarea et al. Evaluating PADM in multiple gingival recession treatments using a modified coronally advanced technique also included Miller Class III recessions [21].

In 2019, Gurlek et al. compared PADM and CTG and combined these materials with a modified coronal advanced flap in the treatment of multiple recessions of Miller class I and II [1, 22]. Both procedures were very effective during the 18-month follow-up, although CTG resulted with a higher gain in keratinized tissue. PADM suffered greater tissue shrinkage and a higher probing depth compared to CTG [23].

In contrast, Pietruska et al. compared the results of both procedures in the treatment of multiple gingival recessions of Miller Class I and II in the mandible via a modified coronal advanced tunnel technique (MCAT) [24, 25]. The mean root coverage data on PADM was 53.20% and 83.10% with the CTG after one year. Complete recession coverage was 20% for PADM and 67% in CTG. The results favour CTG, but these statistical results do not fundamentally influence clinics.

Rakasevic et al. studied 20 patients with multiple adjacent gingival recessions comparing CTG and PADM. They found no significant differences in clinical and aesthetic outcomes using MCAT recorded for 6 and 12 months; however, the mean root coverage was statistically higher in patients treated with CTG. Twice as many patients presented complete root coverage when treated with CTG after 12 months [26].

Similar results between outcomes of multiple gingival recession using CTG and PADM were found in a 20-patient study using a coronal advanced flap (CAF) technique. Increased keratinized tissue, reduction in recession, and complete recession coverage was achieved in both cases. However, as seen in previous studies, more complete recession coverage was obtained in treatments with CTG than PADM especially when keratinized tissue is rare [4, 27].

There are few long-term studies using PADM as a collagen matrix in gingival recession treatment. Cosgarea et al. continued a study initiated in 2016 on 12 patients with a 12-month follow-up mentioned earlier in this article, but extended the follow-up period for four years with a smaller sample of 9 patients. The results demonstrated that the mean recession coverage after 4 years was significantly lower than after 12 months but higher than baseline. Other measured parameters improved after surgery such as recession depth, recession width, width of keratinized tissue and attached gingiva, probing depth, and clinical attachment level [28].

A recent publication by Vincent Bugnas et al. comparing CTG with PADM also using the MCAT technique in 12 patients in a 12 month follow-up proved in this case that CTG obtained better results for mean recession coverage 80.6% ± 23.7% compared to 68.8% ± 23.4% in PADM and also in complete recession coverage 48.7% ± 6.8% with CTG and 24.3% ± 8.2% using PADM. In this study, all parameters that were recorded were in favour of the CTG procedures. However, as we mentioned earlier, they also demonstrated a reduced morbidity and less postoperative pain [29].

The latest publication comparing CTG and PADM this time using the CAF procedure in a study carried out in Brazil on 42 patients divided equally into the control (CTG) group and the test group (PADM), resulting in 18 patients in each group and on a 12 month follow-up, also resulted in better results for CTG specially in MRC in which CTG obtained 91.79 ± 10.1 in comparison with the test group result which was much lower 80.19 ± 16.3. Also, keratinized tissue gain was higher in the control group 0.99 ± 1.23 compared with 0.63 ± 0.83 in the test group [30].

According to recent literature, CTG provides greater gingival margin stability than PADM over time. In a review of 2554 gingival recessions on 1864 patients carried out in 2019, CTG provided a higher recession coverage than PADM and a less tendency for gingival recession recurrence [31].

In summary, gingival recessions have been treated with PADM and showed generally favourable results. However, it is still early to use PADM as a substitute for autogenous connective tissue grafts in mucogingival surgery; more studies are needed to defend this position with longer-term evaluations.
procedures carried out with CTG or PADM. [33, 34].

...levels of plaque deposits around implants and deepening of the vestibule; some buccal KM contraction was observed [39].

Table 1: Porcine acellular dermal matrix in recession treatment: comparing mean recession coverage and keratinized tissue width between procedures carried out with CTG or PADM.

| Authors               | Sample  | Type of study                     | Follow-up time | Surgical procedure | Mean recession coverage (%) | Keratinized tissue width gain (mm) |
|-----------------------|---------|-----------------------------------|----------------|--------------------|------------------------------|-----------------------------------|
| Schmitt et al. [3]    | 8 dogs  | Preclinical study                  | 10 months      | Tunnel technique   | MRC: not registered         | KTW: not registered              |
| Cieslik-Wegemund et al. [2] | 28 patients | Randomized clinical trial | 6 months       | Tunnel technique | CTG: 95% PADM: 91%          |                                   |
| Scuelan et al. [21]   | 12 patients | Case series                       | 12 months      | MCAT               | 73.20 ± 27.71% No CTG control |                                   |
| Gürlek et al. [1]     | 12 patients | Split-mouth randomized clinical trial | 18 months     | CAF                | MRC: not registered         | CTG: 0.51 ± 0.60 PADM: 0.32 ± 0.52 |
| Pietruska et al. [24] | 29 patients | Randomized clinical trial         | 12 months      | MCAT               | CTG: 83.4% PADM: 53.20%     |                                   |
| Rakasevic et al. [26] | 20 patients | Randomized clinical trial         | 12 months      | MCAT               | CTG: 2.96 ± 11.8% PADM: 1.71 ± 13.7% |                                   |
| Maluta et al. [4]     | 15 patients | Split-mouth randomized clinical trial | 6 months  | CAF                | CTG: 95.28 ± 6.89% PADM: 92.68 ± 7.35% |                                   |
| Cosgarea et al. [28]  | 9 patients | Case series                       | 48 months      | MCAT               | PADM: 56.79 ± 27.53% No CTG control |                                   |
| Vincent Bugnas et al. [29] | 12 patients | Random split mouth clinical trial | 12 months | CAF                | PADM: 7.35% PADM: 80.19 ± 6.3% |                                   |
| Meza- Mauricio et al. [30] | 42 patients | Randomized controlled clinical trial | 12 months | CAF                | PADM: 7.35% PADM: 80.19 ± 6.3% |                                   |

3. Comparison with Connective Tissue Graft in Implant Surgery for Soft Tissue Augmentation

Beyond gingival recession, PADM is also used in implant surgery for soft tissue augmentation [32]. It can increase keratinized mucosa (KM) width and thickness. Although there is no official consensus with specific guidelines for the requirement of KM or a minimum volume of peri-implant mucosa required to prevent peri-implant disease, there are many publications clinically proving that the lack of KM will increase the levels of plaque deposits around implants [33, 34].

Recent studies have also demonstrated the importance of having more than 2 mm of KM around implants to reduce the risk of peri-implant disease [35]. Publications by Linkewicz et al. in 2009 showed lower marginal bone loss in implants that had more than 2 mm of KM around them [36]. Other more recent studies proved the need for a minimum of 2 mm of KM to minimize the risk of peri-implantitis [37, 38].

PADM is also being used as an alternative for soft augmentation procedures in implant dentistry substituting the gold standard (CTG). A clinical study published in 2014 by Nocini et al. tested PADM in an extensive keratinized tissue augmentation with deepening of the vestibule. They collected data over different periods: 9 days, 14 days, as well as 1 and 2 months. There was noticeable augmentation of KM around implants and deepening of the vestibule; some buccal KM contraction was observed [39].

Considerable KM gain in width was also observed. One prospective pilot cohort study by Papi et al. used PADM in peri-implant soft tissue augmentation on a second-stage surgery with a small sample of 12 patients. This study did not include a control group using CTG. The KM width increased from 1.35 ± 0.32 mm to 7.86 ± 3.22 mm after a month and 5.67 ± 2.12 mm after a year with an increase of 72.13%. Shrinkage was also observed especially from the first month to the 12th month [40].

Later, the same authors published an article in which a prospective cohort study of a two-year follow-up period in an early implant placement surgery and a concomitant peri-implant augmentation was performed. PADM was combined with synthetic bone in the aesthetic zone. Keratinized mucosa width and gingival thickness were assessed in different periods of 1, 3, 6, 12, and 24 months. Both KM width and gingival thickness increased from baseline to the first month and then decreased from the 1st to the 12th month and remained stable from 12 to 24 months. After 24 months, a 1.94 ± 0.05 mm gain was observed in gingival thickness and a 1.60 ± 0.11 mm gain in KM width compared to baseline [32].

Another important value measured after soft augmentation procedures with PADM as mentioned earlier is soft tissue thickness. As early as 2015, a 6-month follow-up and a sample of 27 patients by Zafiropoulous et al. concluded that PADM leads to a significant increase in soft tissue thickness of 1.06 mm. Although this study had a control group without using any type of graft, previous publications state that the use of CTG increases tissue thickness to an average of 1.2–1.75 mm, which is higher than PADM [41, 42].

Excellent
histological integration and substitution of soft tissue was observed; hence, PADM can be an alternative to CTG [43].

Similar results were observed in soft tissue thickness gain in a longer follow-up of 12 months by Stefanini et al. This study followed 10 patients utilizing the coronal advanced flap surgical technique put together with PADM. There was a 1.2 ± 0.18 mm gain of tissue thickness in the aesthetic zone one year after the final restoration. The authors also found compliance with aesthetic-functional requirements in implant sites. However, this study had no control group [44].

Papi et al. also reported on 12 patients a year later in 2020, where PADM was used for soft tissue augmentation. A second-stage surgery with a 12-month follow-up also demonstrated a gain in tissue thickness this time with a mean gain of 1.25 mm. This is a slightly higher figure (mucosal thickness) than in Stefanini et al. This recent study contrasts with previous ones and also included a three-dimensional volumetric measurement analysis. The first one was published on the buccal contour. The area measured for volume analysis showed a 51.501 mm³ mean gain and 1.25 mm gain of tissue thickness this time with a 3D follow-up and including a test and control group [45].

Schmitt et al. also compared CTG and PADM in this case with a 3D follow-up and including a test and control group with increasing CTG-measured tissue thickness and volume. The volume increase after the six-month period was 19.56 ± 8.95 mm³ for PADM and 61.75 ± 52.69 mm³ for CTG. In the case of tissue thickness in the area of the buccal contour, PADM increased thickness by 0.30 ± 0.16 mm versus CTG 0.80 ± 0.61 mm both after 6 months. The authors concluded that CTG had better results than PADM for both values (soft tissue thickness and volume increase) [46].

A final study, recently published, compared CTG and PADM regeneration techniques to gain keratinized tissue thickness carried out in pigs clinically and histologically examined in three different spans of time: 15 days, 45 days, and 90 days. This showed that the group with PADM had average keratin layer thickness values on top of the CTG group [48]. From 15 to 45 days, there was notable resorption as in previous studies such as Eeckhout et al. and Stefanini et al.

Another important consideration is that CTG when extracted can be of different thicknesses in contrast to PADM which has the same thickness. In further studies, it may be necessary to standardize completely and introduce the same thickness of CTGs for examination.

A summary of significant research studies comparing CTG and PADM in soft tissue augmentation procedures measuring keratinized mucosa and soft tissue thickness is presented in Table 2.

### 4. Conclusion

Randomized clinical trials have been published using strictly porcine acellular dermal matrix as a collagen matrix in dental recession treatment or for soft tissue augmentation in different periods of time: before surgery, immediately after surgery, and after 3 months, 12 months, and 36 months. After surgery, the mean volume increase was 1.53 mm, and after 3 months, it decreased to 1.02 mm due to shrinkage and PADM resorption [40]. After this, a 0.66 mm gain was observed that the authors attribute to the permanent restoration placement pushing the buccal soft tissue [47].

The difference with previous studies such as Stefanini and Papi is that the PADM was placed the same day during the implant surgery. Moreover, in contrast to previous studies like Zafiropoulos and Stefanini which had shorter follow-up periods, these studies carried out measurements with less accurate methods such as transmucosal probing with instruments used in endodontics or anesthetic needles. There were no intermediate measurements after surgery.

Table 2: Soft tissue augmentation in implant sites: comparing keratinized mucosa and soft tissue thickness in procedures using PADM or CTG.

| Authors               | Sample | Study design                                      | Follow-up time | Keratinized mucosa width gain (mm) | Soft tissue thickness gain (mm) |
|-----------------------|--------|---------------------------------------------------|----------------|-----------------------------------|--------------------------------|
| Zafiropoulos et al.   | 27 patients | Prospective, randomized examiner-blind controlled clinical study | 6 months | Not registered | PADM: 1.06 CTG: no CTG control |
| Papi and Pompa        | 12 patients | Prospective pilot cohort study                   | 12 months | PADM: 4.32 | Not registered |
| Stefanini et al.      | 10 patients | Case series                                       | 12 months | 0.65 ± 0.41 | PADM: 1.2 ± 0.18 CTG: no CTG control |
| Papi et al.           | 20 patients | Prospective cohort study                          | 24 months | PADM: 1.60 ± 0.11 | No CTG control |
| Eeckhout et al.       | 15 patients | Prospective case series                           | 36 months | Not registered | PADM: 0.66 CTG: 0.80 ± 0.61 PADM: 0.30 ± 0.16 |
| Schmitt et al.        | 14 patients | Controlled clinical trial                         | 6 months | Not registered | PADM: 1.25 CTG: no CTG control |
| Papi et al.           | 12 patients | Prospective cohort study                          | 12 months | Not registered | PADM: 0.30 ± 1.61 CTG: no CTG control |

Randomized clinical trials have been published using strictly porcine acellular dermal matrix as a collagen matrix in dental recession treatment or for soft tissue augmentation in.
implant surgery. Fewer prospective long-term (>5 years) longitudinal studies exist. The porcine acellular dermal matrix is used in periodontal plastic (gingival recession treatment) and implant surgery and has favourable results as shown in several publications. According to the consensus of the publications reviewed, PADM while providing a benefit in root coverage procedures, it fell short of the outcomes achieved with CTG. At implant sites, soft tissue augmentation procedures offered a favourable outcome similar to CTG for increasing soft tissue thickness.

More studies are needed to compare PADM and CTG using three-dimensional techniques via STL and CBCT superimposition. It is still too early to decide on PADM versus the gold standard (CTG). More evidence is needed including longer trials and larger patient cohorts. PADM has many advantages especially in terms of reduced morbidity and reduced surgical times.

Data Availability

All data in the article are cited in references.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

References

[1] Ö. Gürlek, P. Gümüş, N. Nizam, and N. Buduneli, “Coronally advanced flap with connective tissue graft or xenogeneic acellular dermal matrix in the treatment of multiple gingival recessions: a split-mouth randomized clinical trial,” Journal of Esthetic and Restorative Dentistry, vol. 32, no. 4, pp. 380–388, 2020.

[2] M. Cie´ slik-Wegemund, B. Wierucka-Młynarczyk, M. Tanasiewicz, and Ł. Gilowski, “Tunnel technique with collagen matrix compared with connective tissue graft for treatment of periodontal recession: a randomized clinical trial,” Journal of Periodontology, vol. 87, no. 12, pp. 1436–1443, 2016.

[3] C. M. Schmitt, R. E. Matta, T. Moest et al., “Soft tissue volume alterations after connective tissue grafting at teeth: the subepithelial autologous connective tissue graft versus a porcine collagen matrix - a pre-clinical volumetric analysis,” Journal of Clinical Periodontology, vol. 43, no. 7, pp. 609–617, 2016.

[4] R. Maluta, M. F. Monteiro, D. C. Peruzzo, and J. C. Joly, “Root coverage of multiple gingival recessions treated with coronally advanced flap associated with xenogeneic acellular dermal matrix or connective tissue graft: a 6-month split-mouth controlled and randomized clinical trial,” Clinical Oral Investigations, vol. 9, 2021.

[5] J. R. Wessel and D. N. Tatakis, “Patient outcomes following subepithelial connective tissue graft and free gingival graft procedures,” Journal of Periodontology, vol. 79, no. 3, pp. 425–430, 2008.

[6] M. Del Pizzo, F. Modica, N. Bethaz, P. Priotto, and R. Romagnoli, “The connective tissue graft: a comparative clinical evaluation of wound healing at the palatal donor site. a preliminary study,” Journal of Clinical Periodontology, vol. 29, no. 9, pp. 848–854, 2002.

[7] R. J. Harris, R. Miller, L. H. Miller, and C. Harris, “Complications with surgical procedures utilizing connective tissue grafts: a follow-up of 500 consecutively treated cases,” The International Journal of Periodontics and Restorative Dentistry, vol. 25, no. 5, pp. 449–459, 2005.

[8] R. Burkhardt, C. H. Hämmerle, and N. P. Lang, “Research group on oral soft tissue biology & wound healing. self-reported pain perception of patients after mucosal graft harvesting in the palatal area,” Journal of Clinical Periodontology, vol. 42, no. 3, pp. 281–287, 2015.

[9] L. A. Aguirre-Zorrano, A. M. García-De La Fuente, R. Estefanía-Fresco, and X. Marichalar-Mendía, “Complications of harvesting a connective tissue graft from the palate. a retrospective study and description of a new technique,” Journal of Clinical and Experimental Dentistry, vol. 9, no. 12, pp. e1439–e1445, 2017.

[10] D. J. Wainwright, “Use of an acellular allograft dermal matrix (AlloDerm) in the management of full-thickness burns,” Burns, vol. 21, no. 4, pp. 243–248, 1995.

[11] J. Aragoneses, A. Suárez, C. Rodríguez, and J. M. Aragoneses, “Histomorphometric comparison between two types of acellular dermal matrix grafts: a mini pig animal model study,” International Journal of Environmental Research and Public Health, vol. 18, no. 8, p. 3881, 2021.

[12] D. R. B. de Resende, S. L. A. Greghi, A. F. Siqueira, C. A. M. Benfatti, C. A. Damante, and M. S. Ragghianti Zangrando, “Acellular dermal matrix allograft versus free gingival graft: a histological evaluation and split-mouth randomized clinical trial,” Clinical Oral Investigations, vol. 23, no. 2, pp. 539–550, 2019.

[13] R. J. Harris, “Root coverage with a connective tissue with partial thickness double pedicle graft and an acellular dermal matrix graft: a clinical and histological evaluation of a case report,” Journal of Periodontology, vol. 69, no. 11, pp. 1305–1311, 1998.

[14] M. E. Aichelmann-Reidy and R. A. Yukna, “Bone replacement grafts. The bone substitutes,” Dental Clinics of North America, vol. 42, no. 3, pp. 491–503, 1998.

[15] D. Rothamel, M. Benner, T. Fienitz et al., “Biodegradation pattern and tissue integration of native and cross-linked porcine collagen soft tissue augmentation matrices - an experimental study in the rat,” Head & Face Medicine, vol. 10, p. 10, 2014.

[16] A. M. Pabst, A. Happe, A. Callaway et al., “In vitro and in vivo characterization of porcine acellular dermal matrix for gingival augmentation procedures,” Journal of Periodontal Research, vol. 49, no. 3, pp. 371–381, 2014.

[17] P. Papi and G. Pompa, “The use of a novel porcine derived acellular dermal matrix (macoderm) in peri-implant soft tissue augmentation: preliminary results of a prospective pilot cohort study,” BioMed Research International, vol. 2018, Article ID 6406051, 2018.

[18] A. M. Pabst, K. M. Lehmann, C. Walter, M. Krüger, S. I. Stratul, and A. Kasaj, “Influence of porcine-derived collagen matrix on endothelial progenitor cells: an in vitro study,” Odontology, vol. 104, no. 1, pp. 19–26, 2016.

[19] M. A. Saghiri, A. Asatourian, F. Garcia-Godoy, and N. Sheibani, “The role of angiogenesis in implant dentistry part II: the effect of bone-grafting and barrier membrane materials on angiogenesis,” Medicina Oral, Patología Oral y Cirugía Bucal, vol. 21, no. 4, pp. e526–e537, 2016.

[20] Z. Lin, C. Nica, A. Sculean, and M. B. Asparuhova, “Enhanced wound healing potential of primary human oral fibroblasts and periodontal ligament cells cultured on four different porcine-derived collagen matrices,” Materials, vol. 13, no. 17, p. 3819, 2020.
[21] A. Sculean, R. Cosgarea, A. Stähli et al., “Treatment of multiple adjacent maxillary Miller Class I, II, and III gingival recessions with the modified coronally advanced tunnel, enamel matrix derivative, and subepithelial connective tissue graft: a report of 12 cases,” *Quintessence International*, vol. 47, no. 8, pp. 653–659, 2016.

[22] G. Zucchelli and M. De Sanctis, “Treatment of multiple recession-type defects in patients with aesthetic demands,” *Journal of Periodontology*, vol. 71, no. 9, pp. 1506–1514, 2000.

[23] F. Vignoletti, J. Nuñez, N. Discopoli et al., “Clinical and histological healing of a new collagen matrix in combination with the coronally advanced flap for the treatment of Miller class-I recession defects: an experimental study in the minipig,” *Journal of Clinical Periodontology*, vol. 38, no. 9, pp. 847–855, 2011.

[24] M. Pietruska, A. Skurska, Ł. Podlewski, R. Milewski, and J. Pietruski, “Clinical evaluation of Miller class I and II recessions treatment with the use of modified coronally advanced tunnel technique with either collagen matrix or subepithelial connective tissue graft: a randomized clinical study,” *Journal of Clinical Periodontology*, vol. 46, no. 1, pp. 86–95, 2019.

[25] S. Parween, J. P. George, and M. Prabhuj, “Treatment of multiple mandibular gingival recession defects using MCAT technique and SCTG with and without RHPDGF-BB: a randomized controlled clinical trial,” *The International Journal of Periodontics and Restorative Dentistry*, vol. 40, no. 2, pp. e43–e51, 2020.

[26] D. L. Rakasevic, I. Z. Milinkovic, S. M. Jankovic, I. A. Soldatovic, Z. M. Aleksi, and N. S. Nikolic-Jakoba, “The use of collagen porcine dermal matrix and connective tissue graft with modified coronally advanced tunnel technique in the treatment of multiple adjacent type I gingival recessions: a randomized, controlled clinical trial,” *Journal of Esthetic and Restorative Dentistry*, vol. 32, no. 7, pp. 681–690, 2020.

[27] M. Del Pizzo, G. Zucchelli, F. Modica, R. Villa, and C. Debernardi, “Coronally advanced flap with or without enamel matrix derivative for root coverage: a 2-year study,” *Journal of Clinical Periodontology*, vol. 32, no. 11, pp. 1181–1187, 2005.

[28] R. Cosgarea, R. Miron, R. Bora, A. Rosu, S. Buduru, and A. Sculean, “Long-term results after treatment of multiple adjacent gingival recessions with the modified coronally advanced tunnel and a porcine acellular dermal matrix,” *Quintessence International*, vol. 52, no. 1, pp. 32–44, 2021.

[29] S. Vincent-Bugnas, J. Laurent, E. Namay, M. Charbit, and G. Borie, “Treatment of multiple gingival recessions with xenogeneic acellular dermal matrix compared to connective tissue graft: a randomized split-mouth clinical trial,” *Journal of Periodontal & Implant Science*, vol. 35, no. 3, pp. 77–87, 2021.

[30] J. Meza-Mauricio, J. Cortez-Gianezzi, P. M. Duarte, L. Tavelli, G. Rasperini, and M. de Faveri, “Comparison between a xenogeneic dermal matrix and connective tissue graft for the treatment of multiple adjacent gingival recessions: a randomized controlled clinical trial,” *Clinical Oral Investigations*, vol. 82, 2021.

[31] L. Tavelli, S. Barootchi, F. Cairo, G. Rasperini, K. Sheddon, and H. L. Wang, “The effect of time on root coverage outcomes: a network meta-analysis,” *Journal of Dental Research*, vol. 98, no. 11, pp. 1195–1203, 2019.

[32] P. Papi, N. Pranno, B. Di Murro, and G. Pompa, “Early implant placement and peri-implant augmentation with a porcine-derived acellular dermal matrix and synthetic bone in the aesthetic area: a 2-year follow-up prospective cohort study,” *International Journal of Oral and Maxillofacial Surgery*, vol. 50, no. 2, pp. 258–266, 2021.

[33] D. S. Thoma, B. Buranawat, C. H. Hämmerle, U. Held, and R. E. Jung, “Efficacy of soft tissue augmentation around dental implants and in partially edentulous areas: a systematic review,” *Journal of Clinical Periodontology*, vol. 41, no. 15, pp. S77–S91, 2014.

[34] D. M. Chung, T. J. Oh, J. L. Shotwell, C. E. Misch, and H. L. Wang, “Significance of keratinized mucosa in maintenance of dental implants with different surfaces,” *Journal of Periodontology*, vol. 77, no. 8, pp. 1410–1420, 2006.

[35] A. Monje and G. Blasi, “Significance of keratinized mucosa/gingiva on peri-implant and adjacent periodontal conditions in erratic maintenance compliers,” *Journal of Periodontology*, vol. 90, no. 5, pp. 445–453, 2019.

[36] T. Linkevicius, P. Apse, S. Grybauskas, and A. Puisys, “The influence of soft tissue thickness on crestal bone changes around implants: a 1-year prospective controlled clinical trial,” *The International Journal of Oral & Maxillofacial Implants*, vol. 24, no. 4, pp. 712–719, 2009.

[37] A. S. Gharpure, J. M. Latimer, F. E. Alijof, and D. M. Daubert, “Role of thin gingival phenotype and inadequate keratinized mucosa width (< 2 mm) as risk indicators for peri-implantitis and peri-implant mucositis,” *Journal of Periodontology*, vol. 15, 2021.

[38] V. Moraschini, D. Luz, G. Velloso, and E. D. P. Barboza, “Quality assessment of systematic reviews of the significance of keratinized mucosa on implant health,” *International Journal of Oral and Maxillofacial Surgery*, vol. 46, no. 6, pp. 774–781, 2017.

[39] P. F. Nocini, R. Castellani, G. Zanotti et al., “Extensive keratinized tissue augmentation during implant rehabilitation after Le Fort I osteotomy: using a new porcine collagen membrane (Mucoderm),” *Journal of Craniofacial Surgery*, vol. 25, no. 3, pp. 799–803, 2014.

[40] F. Vignoletti, J. Nuñez, F. de Sanctis, M. Lopez, R. Caffesse, and M. Sanz, “Healing of a xenogeneic collagen matrix for keratinized tissue augmentation,” *Clinical Oral Implants Research*, vol. 26, no. 5, pp. 545–552, 2015.

[41] F. Cairo, L. Barbato, P. Tonelli, G. Batalocco, G. Pagavino, and M. Nieri, “Xenogeneic collagen matrix versus connective tissue graft for buccal soft tissue augmentation at implant site. a randomized, controlled clinical trial,” *Journal of Clinical Periodontology*, vol. 44, no. 7, pp. 769–776, 2017.

[42] J. Gargallo-Albiol, S. Barootchi, L. Tavelli, and H. L. Wang, “Efficacy of xenogeneic collagen matrix to augment peri-implant soft tissue thickness compared to autogenous connective tissue graft: a systematic review and meta-analysis,” *The International Journal of Oral & Maxillofacial Implants*, vol. 34, no. 5, pp. 1059–1069, 2019.

[43] G. G. Zafiropoulos, G. Deli, O. Hoffmann, and G. John, “Changes of the peri-implant soft tissue thickness after grafting with a collagen matrix,” *Journal of Indian Society of Periodontology*, vol. 20, no. 4, pp. 441–445, 2016.

[44] M. Stefanini, A. Rendon, and G. Zucchelli, “Porcine-derived acellular dermal matrix for buccal soft tissue augmentation at single implant sites: a 1-year follow-up case series,” *The International Journal of Periodontics and Restorative Dentistry*, vol. 40, no. 1, pp. 121–128, 2020.

[45] P. Papi, D. Penna, B. Di Murro, and G. Pompe, “Clinical and volumetric analysis of peri-implant soft tissue augmentation using an acellular dermal matrix: a prospective cohort study,” *Journal of Periodontology*, vol. 19, 2020.
[46] C. M. Schmitt, P. Brückbauer, K. A. Schlegel, M. Buchbender, W. Adler, and R. E. Matta, "Volumetric soft tissue alterations in the early healing phase after peri-implant soft tissue contour augmentation with a porcine collagen matrix versus the autologous connective tissue graft: a controlled clinical trial," *Journal of Clinical Periodontology*, vol. 48, no. 1, pp. 145–162, 2021.

[47] C. Eeckhout, E. Bouckaert, D. Verleyen, T. De Bruyckere, and J. Cosyn, "A 3-year prospective study on a porcine-derived acellular collagen matrix to re-establish convexity at the buccal aspect of single implants in the molar area: a volumetric analysis," *Journal of Clinical Medicine*, vol. 9, no. 5, p. 1568, 2020.

[48] J. Aragoneses, A. Suárez, C. Rodríguez, and J. M. Aragoneses, "Clinical and histological differences between guided tissue regeneration with acellular dermal matrix of porcine origin and autologous connective tissue: an animal study," *Materials*, vol. 14, no. 2, p. 272, 2021.