

Periodontal regenerative medicine using mesenchymal stem cells and biomaterials: A systematic review of pre-clinical studies

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The aim of the systematic review was to analyze the use of mesenchymal stem cells (MSC) and biomaterial for periodontal regeneration from preclinical animal models and human. Electronic databases were searched and additional hand-search in leading journals was performed. The research strategy was achieved according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines. The including criteria were as follows: MSC, biomaterial, *in vivo* studies, with histologic and radiologic analysis and written in English. The risk of bias was assessed for individual studies. A total of 50 articles were selected and investigated in the systematic review. These results indicate that MSC and scaffold provide beneficial effects on periodontal regeneration, with no adverse effects of such interventions. Future studies need to identify the suitable association of MSC and biomaterial and to characterize the type of new cementum and the organization of the periodontal ligament fiber regeneration.

Keywords: Mesenchymal stem cells, Biomaterial, Periodontal regeneration, Pre-clinical study

INTRODUCTION

The periodontium is a highly specialized and dynamic tissue. It consists in a tooth-anchoring device made of two soft tissues, the gingiva and the periodontal ligament, and of two hard tissues, the cement and the alveolar bone. The gingiva, part of the superficial periodontium, is composed of an epithelium and a connective tissue, forming a periodontal attachment system that allows fluid exchange and assures a complete crimp around the tooth¹. As for the deep periodontium, made of the periodontal ligament, the cement and the alveolar bone, it acts like an alveolar anchorage system, enabling both stability and damping of the tooth. In the oral environment, the periodontium is confronted with more than 1,000 bacteria species². Thus, the space between the tooth and the surrounding gingiva, called sulcus, can become a gateway to potential inflammatory diseases. The most common cause of periodontal destruction is periodontitis^{3,4}. This multifactorial disease is due to an opportunistic bacteria contamination on a specific site, paired with local risk factors (e.g., insufficient oral hygiene, decay) and general risk factors (e.g., HIV, diabetes). In response to this bacterial aggression, an inflammatory process in the gingival sulcus will slowly destroy the periodontal structures and attachment, leading to tooth mobility and premature tooth loss⁵.

Conventional treatment strategies are based on a sustained decrease in the microbial load through a non-surgical or a surgical elimination of the dental plaque associated with assiduous plaque control from the patient. In spite of great progress in the understanding

of the pathogenesis of periodontitis, the tools to treat it seem to only postpone the unavoidable tooth loss linked to periodontal disease, and fail to restore *ad integrum* periodontal tissue, proving it unsatisfactory both for patients and for dental surgeons. Since the 1970s, several procedures have been attempted to restore such lost tissues, including autogenous bone grafting, implantation of biomaterials including bone derivatives and bone substitutes, guided-tissue regeneration (GTR) procedures⁶, and implantation of biologic factors, including enamel matrix proteins^{7,8}. Still, these strategies fail to regenerate the complete periodontium damage since the quality of repaired tissue remain variable and limited^{9,10}. To date, complete periodontal regeneration is not achievable in a highly reproducible and easy way. Therefore, functional and aesthetic sequelae are commonly found in treated patients with a history of periodontitis.

That is why, re-establishing the original structure, properties and functions of the diseased periodontium remains a significant clinical challenge. To address this issue, regenerative medicine using an effective combination of mesenchymal stem cells (MSC) and biomaterial have become subjects of particular interest¹¹. Thus, a wide variety of the studies focus on MSC and try to combine their potential with suitable biomaterials in order to obtain a periodontal regeneration¹². Despite numerous publications in pre-clinical animal models and humans, the efficacy of the association of MSC and biomaterial for periodontal regeneration remain controversial^{13,14}.

To increase the value of individual preclinical studies as proof of concept for randomized clinical trials, systematic reviews have been proposed as the

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standard method for analyzing experimental work involving animals¹⁵⁾. Systematic review is an essential tool for summarizing evidence reliably and accurately. It provides a starting point for guideline developers for clinical practice. In the case of regenerative medicine for the enhancement of periodontal regeneration, a systematic review may provide valuable information for a suitable clinical practice.

In this context, the purpose of the present systematic review was to assess the scientific literature to obtain more clarity on the efficacy of periodontal regeneration strategies using the association of MSC and biomaterials in pre-clinical animal and human studies.

MATERIALS AND METHODS

The different studies concerning periodontal regeneration with the help of MSC and biomaterials on human or animal models have been collected and analyzed.

Information sources and search strategy

The research strategy was achieved according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines¹⁵⁾ as much as possible. Original articles were searched using electronic and manual databases until March 2018. Furthermore, relevant articles were screened by hand to potentially add relevant new articles.

This search was applied to Medline, Cochrane Library, Lilacs and Report Evidence-based Practice Centers (EPC). Electronic searches were completed by additional hand-search performed for the International Journal of Periodontics and Restorative Dentistry, the Journal of Clinical Periodontology, the Journal of Dental Research, the Journal of Periodontal Research and the Journal of Periodontology.

The following Medical Subject Heading (MeSH) terms and keywords were used: "stem cell", "periodontal", "regeneration", "biomaterial", and "*in vivo*". Only English articles were included, and no publication dates or publication status restrictions were imposed.

Study selection and inclusion/exclusion criteria

For the selection of studies, two investigators (SP, XS) screened the titles and the abstracts of the publications in an unblended, standardized manner. Selection was

based on the inclusion and exclusion criteria defined so as to include only the most valuable articles (Table 1). Studies deemed to meet the inclusion criteria and those with insufficient information to make a clear decision were selected. The second phase consisted of assessing the whole articles by the same investigators to determine the eligibility of the study. The selection process was recorded in detail to a PRISMA flow diagram (Fig. 1). Any disagreements between the two investigators regarding inclusion of a study were resolved by discussion.

Data collection process and data items

The characteristics of the study were extracted independently by the same investigators and recorded. The data were compared for accuracy and any discrepancies were discussed and resolved by consensus.

Both reviewers extracted from the included studies the following data: (1) Cell type, passage number, differentiation, number per defect; (2) Biomaterial (+/-membrane); (3) Animal models: species, strain, sex, age, weight; (4) Number of defects per group; (5) Defect type, size, induced inflammation; (6) Treatment groups; (7) Observation period; (8) Qualification of newly formed tissues; (9) Results (Table 2). If one of these data is not reported in the Table, it means that the information is not mentioned by the authors.

Risk of bias in individual studies

To ascertain the risk of bias in eligible articles, the same investigators in a blind manner evaluated their methodology either by SYRCLE's Risk of Bias tool for animal intervention studies¹⁶⁾, or by Risk Of Bias in Nonrandomized Studies of Interventions (ROBINS-I) tool¹⁷⁾ modified for human non-randomized trials; or by the Cochrane Collaboration's tool for human randomized trials¹⁸⁾. The case reports were excluded from the risk of bias analysis because of the absence of adapted methodology.

Regarding SYRCLE's Risk of Bias tool, the unit of analysis errors is assessed as a "high risk of bias" if the interventions to parts of the body within one participant (*i.e.*, splits mouth: control and experimental side) was reported. In addition, many items remain as "unclear" due to the poor description of methodology. In order to complete the investigation, two items were added

Table 1 Inclusion and exclusion criteria

Inclusion criteria:	Exclusion criteria:
✓ Studies using MSC	- Studies using only MSC
✓ Studies using biomaterial	- Studies using growth factors only
✓ Studies with the precise number of specimens	- <i>in vitro</i> studies
✓ <i>in vivo</i> studies	- <i>in vivo</i> studies on rodents and lagomorphs
✓ Studies with histologic and radiologic analysis	- Studies on implant or bone regeneration only
✓ Studies written in English	- Studies with $p>0.05$
	- Review
	- Studies without control groups

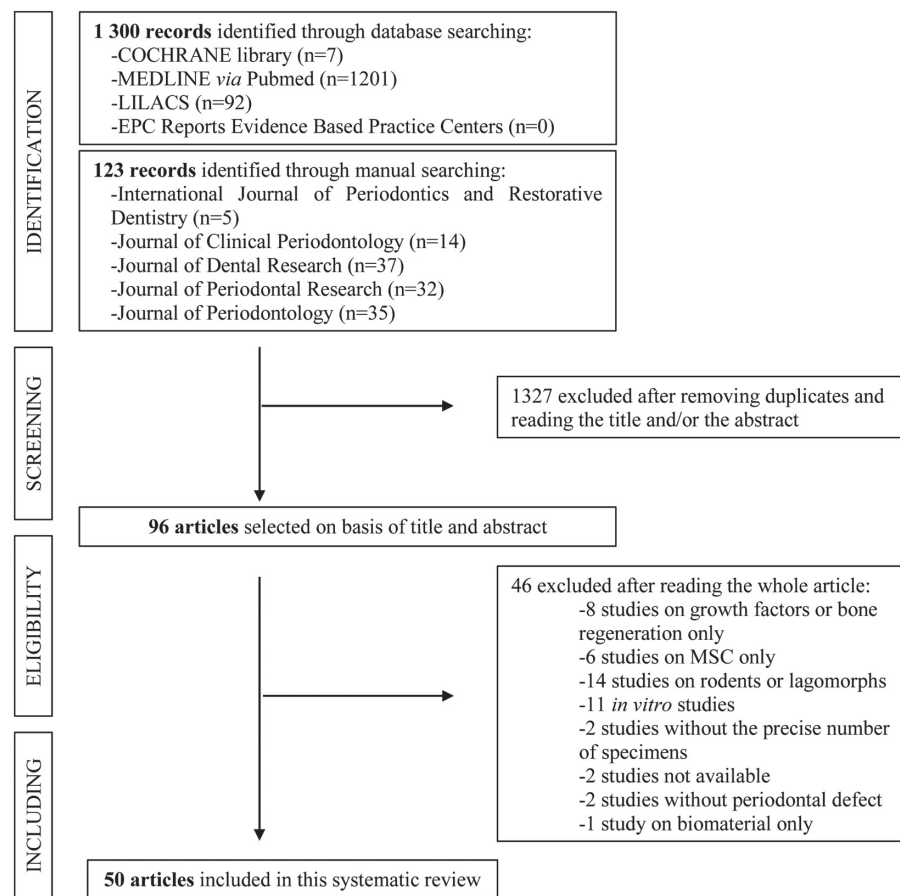


Fig. 1 PRISMA flowchart for identifying eligible studies.

mentioned as follows: "mention of randomization" and "mention of binding".

Concerning the Cochrane Collaboration's and SYRCLE's Risk of Bias tool, we considered that there was no exclusion of animals when the number of animals reported in the method section equaled to the number mentioned in the results section. Disagreements between the investigators were resolved by consensus.

Data synthesis

A meta-analysis could not be performed because of the studies' heterogeneity. Consequently, we conducted a descriptive and systematic analysis of the studies.

RESULTS

Study selection

Taking into account the previously defined criteria, 1,423 studies were initially identified (Fig. 1). The electronic search of Medline, Cochrane library, Lilacs, EPC databases provided 1,300 articles. Additionally, 123 studies were selected by hand-searching the International Journal of Periodontics and Restorative Dentistry, the Journal of Clinical Periodontology, the Journal of Dental Research, the Journal of Periodontal

Research and the Journal of Periodontology. After adjusting for duplicates and reading the title and/or the abstract, 96 studies remained. Out of these, 96 were discarded because after reviewing the whole article, it appeared that these papers clearly did not meet the inclusion criteria. A total of 50 articles were included in the systematic review.

Study characteristics

The studies were then ranked in a comparative Table (Tables 2 and 3) according to the alphabetical order. These tables show a wide variety of combinations of MSC and biomaterials. The review began with an analysis of the type of population, mainly of preclinical animal species. Indeed, experimental models in majority are found in dogs, employed in 25 studies. Twelve studies were performed with miniature pigs and two with sheeps. Only 11 studies have been performed with humans. A variety of periodontal defects were used in the selected studies, including class III furcation defects (7 studies), fenestration defects (3 studies), intrabony with class II furcation defects (1 study), class II furcation defects (6 studies), dehiscence defects (2 studies), a root-shaped implant sockets (2 studies), alveolar sockets (2 studies), combined periodontal-endodontic lesions (1 study) but

Table 2 Comparative table of pre-clinical animal studies in periodontal regeneration combining MSC and biomaterials

Reference	Cell type/ passage number/ differentiation/ number per defect	Biomaterial/+/- membrane	Animal model/ species/sex/age/ weight	Number of defects per group	Defect type/size/ induced inflammation	Treatment groups	Observation period	Qualification of newly formed tissues	Results
Akinci T <i>et al.</i> 2005 ³⁴⁾	-PDLC -P1-6 ·vit C stimulation ·1×10 ⁶ /sheet	Hyaluronic acid sheet	DOG Beagle, female, 3 years-old, 9.8 and 11.2 kg	5 (a, b)	-dehiscence defects -5×5mm ² -no inflammation	(a) PDLC+hyaluronic acid sheet (b) hyaluronic acid sheet	2 months	-Histological: HES, Masson trichrome staining -Histomorphometric	No sign of inflammation or recession in the 2 groups. Formation of cement, bone, and ligament more significant in the group (a) than in the group (b)
Chen G <i>et al.</i> 2015 ³⁵⁾	-DFSC ·undifferentiated	Gelatin electrospun sheet (APES)+treated dentin matrix (TDM)+native dental pulp extracellular matrix (DPEM)	MINIATURE PIG 1 year old	3 (a, b)	-root-shaped implant socket -no inflammation	(a) DFSC+DPEM+APES+TDM (b) TDM	3 months	-Histological -Radiological: micro-CT -Immunohistochemical: Col-1, Col-3, periostin, DMP-1 sialophosphoprotein, GFP	The sandwich composites APES/TDM/DPEM and DFSC generated periodontal ligament-like tissues
Ding G <i>et al.</i> 2010 ³⁶⁾	-PDLSC -P0 ·undifferentiated ·2 cell sheets	HA/β-TCP +gelatin membrane	MINIATURE PIG female Wuzhishan, male Guizhou, 6-8 months old, 30-40 kg	6 (a, b, c, d, e)	-intrabony defects -3×7×5mm ³ -inflammation	(a) PDLSC autologous+HA-β-TCP (b) PDLSC allogenic+HA-β-TCP (c) PDLC autologous and allogenic+HA-β-TCP (d) HA+β-TCP (e) empty defect	3 months	-Clinical: CAL, PD, GR, blood and biochemical tests -Histological: HES -Histomorphometric -Radiological: CT	Allogenic and autogenous PDLSC have improved periodontal regeneration in the same way compared to other groups; with a nearly complete recovery of bone cement and ligament
Dogan A <i>et al.</i> 2002 ³⁶⁾	-regenerated periodontal ligament derived cells -P4 ·undifferentiated ·2×10 ⁶	Autologous blood coagulum	Dog	2 (a, b)	-class II furcation defects -5×2 mm ² -no inflammation	(a) regenerated periodontal ligament derived cells+autologous blood coagulum (b) empty	1, 5 months	-Histological: HES -Histomorphometric	Seeding of cells into periodontal defects promote the bone regeneration and adaptation of connective tissue fibers on exposed dentin surface with limited cementum formation
Dogan A <i>et al.</i> 2003 ³⁷⁾	-regenerated periodontal ligament derived cells -P4 ·undifferentiated ·2×10 ⁶	Autologous blood coagulum	Dog	2 (a, b)	-fenestration defects -5×5 mm ² -no inflammation	(a) regenerated periodontal ligament derived cells+autologous blood coagulum (b) empty	1, 5 months	-Histological: HES -Histomorphometric	In the cell seeding group, the main periodontal healing pattern was connective tissue adaptation, characterized by parallel bundles of collagen fibrils resting on root dentin

Table 2 continued

		MINIATURE PIG 1 female and 7 males, 18±1 months old, 46.9±4.6 kg	(a) GMSC+Bio oss® (b) GMSC+collagen scaffold (c) bio oss® (d) collagen scaffold (e) mucoperiosteal flap and root planning only (f) without intervention	-intradony defects -3×7×5 mm³ -inflammation	3 months	-Clinical: CAL, PD, GR, BOP, PI -Histological: HES -Histomorphometric -Radiological: CT	The treatments with GMSC compared with scaffolds alone or control groups have entailed better results
Fawy El-Sayed KM <i>et al.</i> 2012 ⁽⁴⁾	-GMSC -P3 -undifferentiated -2×10 ⁷	Bio oss® +Bio-Gide®	(a, b, c, d, e, f)	(a, b, c, d) -intrabony defects -3×7×5 mm³ -inflammation	3 months	-Clinical: CAL, PD, GR, BOP, PI -Histological: HES -Histomorphometric -Radiological: CT	
Fawy El-Sayed KM <i>et al.</i> 2015 ⁽⁵⁾	-GMSC -P3 -undifferentiated -2×10 ⁷	IL-1ra-releasing HA-ECM +Bio-Gide®	(a, b, c, d)	(a, b, c, d) -intrabony defects -3×7×5 mm³ -inflammation	4 months	-Clinical: CAL, PD, GR, BOP, PI -Histological: HES -Histomorphometric -Radiological: CT	
Fu X <i>et al.</i> 2014 ⁽⁸⁾	-PDLC or SHED -P3 -vit C stimulation -2 cell sheets	HAβ-TCP	(a, b, c)	(a, b, c, d) -intrabony defects -5×7×7 mm³ -inflammation	3 months	-Clinical: CAL, PD, GR, BOP, PI -Histological: HES -Radiological: CT	No significant difference between the 2 groups PDLC and SHED, both have significantly improved periodontal regeneration with a very small inflammatory area
Gao ZH <i>et al.</i> 2016 ⁽⁹⁾	-PDLC; -P2-P3 -undifferentiated -2×10 ⁶	HA/TCP	(a)	(a) HA+TCP+DPSC+PDLC (b) dental implants	6, 12 months	-Clinical: PD, GR, gingivitis, peri-implantitis -Histological: HES, toluidine blue staining -Radiological: CT, micro-CT -Biochemical: compressive strength, modulus of elasticity, torsional force -SEM	These data showed that the group (a) could function as well as the group (b). Histological staining showed that PDLC like and bone tissue had been generated similar to the natural sample in both groups. Neither the group (a) nor the group (b) showed gingivitis or peri-implantitis. However, the restoration success rate was significantly lower in with the group (a) than with the group (b)
Hasegawa N <i>et al.</i> 2006 ⁽²²⁾	-BMSC -undifferentiated -2×10 ⁷	Atelocollagen	46 (a)	-root-shaped implant socket -4.1×10 mm² -no inflammation	(a) BMSC+ratelocollagen (b) no treatment	-Histological: HES -Immunohistochemical: GFP, Proliferating Cell Nuclear Antigen (PCNA)	Outcome showed that the defects were almost regenerated with cementum, periodontal ligament, and alveolar bone after MSC transplantation
Inukai T <i>et al.</i> 2013 ⁽⁵⁾	-PDLC -undifferentiated -2×10 ⁶	Atelocollagen	5 (b)	-intrabony defects -4×5 mm² -no inflammation	(a) PDLC+ratelocollagen (b) atelocollagen+PBS (c) no treatment	-Histological: HES -Histomorphometric -Radiological: dental X-ray	Both cement and bone heights as well as bone surface were more significant in the group with PDLC
Iwata T <i>et al.</i> 2009 ⁽⁶⁾	-PDLC -P3 -osteoinductive medium -9×10 ⁴	PGAβ-TCP	4 (a, b)	-intrabony defects -5×5×4 mm³ -no inflammation	(a) PDLC+PGAβ-TCP (b) PGAβ-TCP	1, 5 months -Histological: HES, Azan staining -Histomorphometric -Radiological: micro-CT	PDLC permitted bone and cement regeneration with correctly orientated collagen while a limited formation bone was noted in the control group

Table 2 continued

Jiang J et al. 2010 ⁽³⁰⁾	-Periodontal cells -P3 -undifferentiated -5x10 ⁶	β -TCP +ePTFE membranes	DOG Beagle, 9.5 and 10.5 kg (a, b, c)	-class III furcation defects -3x4 mm ² -inflammation	(a) PDLSC+atocollagen (b) atocollagen (c) empty	3 months	-Histological: HES, Mallory's trichrome stain -Histomorphometric -Radiological	PDLSC associated with atocollagen significantly improve periodontal and bone formations compared to control groups (b, c)
Kawaguchi H et al. 2004 ⁽²⁹⁾	-BMSC -P3 -undifferentiated -2x10 ⁶ , 5x10 ⁶ , 1x10 ⁷ , 2x10 ⁷	Atelocollagen	DOG Beagle female, 10-14 kg, 12-20 months old	-class III furcation defects -4 mm in depth -no inflammation	(a) BMSC+atelocollagen (b) atelocollagen	1 month	-Histological: HES, Azan staining -Histomorphometric	In the BMSC group, neocement covered nearly all denuded dentin surface, added to bone and ligament formation (Sharpey's fibers like). But no <i>ad-integrum</i> bone recovery. No root system ankylosis or root resorption
DPSC	-P3 -undifferentiated -2x10 ⁷ /default	3-4 Bio-Oss [®] granules	DOG Mongrel male 1-2 years old, 14-22 kg	-intrabony defects -3x5x8 mm ³ -inflammation	(a) DPSC+Bio-Oss [®] (b) Bio-Oss [®]	2 months	-Histologic: HES -Histomorphometric	The cement generated in the presence of DPSC was thicker and covered a bigger surface compared with the control group. Likewise the formation of ligament was more significant in the test group. However, no noticeable difference in bone formation between the 2 groups
Khorsand A et al. 2013 ⁽³¹⁾	-P3 -undifferentiated -2x10 ⁷ /default							
Lang H et al. 1998 ⁽²⁾	-Primary cell cultures from alveolar bone and periodontal ligament -vit C stimulation -2x10 ⁶ /mL	Gelita [®] +ePTFE membranes	MINIATURE PIG Trol types, 4-8 years old	-intrabony and class II furcation defects -inflammation	(a) cells+Gelita [®] (b) Gelita [®] (c) empty with membrane (d) empty without membrane (e) no treatment	10, 30, 90 days	-Clinical: PI, PD, GR, Bling Index -Histological: polyfluorochrome labeling, toluidine blue staining	The study shows that the primary cell cultures from alveolar bone and periodontal ligament leads to formation of new cementum and bone
Li H et al. 2009 ⁽³²⁾	-Cryopreserved or No-cryopreserved BMSC -P4 -undifferentiated -5x10 ⁶ /mL	Collagen membrane +ePTFE membranes	DOG Beagle, female, 12-18 kg (c)	-penetration defects -5x5 mm ² -no inflammation	(a) BMSC cryopreserved+ collagen (b) BMSC no cryopreserved+ collagen (c) collagen	2 months	-Histological: HES, Masson's staining -Histomorphometric	BMSC whether cryopreserved or not showed a better periodontal regeneration with bone, cement and ligament neformation compared with the group with collagen only
Liu Y et al. 2008 ⁽²⁾	-PDLSC -P3 -undifferentiated -2x10 ⁷	HA β -TCP +gelatin membranes	MINIATURE PIG 12 months old, 30-40 kg (a, b, c)	-intrabony defects -7x3x5 m ³ -inflammation	(a) PDLSC+HA β -TCP (b) HA β -TCP (c) no treatment	3 months	-Clinical: CAL, PI, PD, GR, BOP -Histological: HES, GFP -Radiological: CT	The treatment with PDLSC entailed bone, ligament and cement regeneration with a good anchoring of Sharpey's fibers compared with the other groups
Liu Z et al. 2016 ⁽³³⁾	-BMSC -undifferentiated	collagen- hydroxyapatite scaffold	DOG Beagles, male, 12 months old 10.5-12 kg	-intrabony defects -3x5mm ² -no inflammation	(a) BMSC+HA+collagen (b) BMSC+HA+collagen-cross linked (c) empty (d) no treatment	3 dogs: 3 months 3 dogs: 6 months	-Histological: HES, Masson's staining -Radiological: Micro-CT	In group (a, b), newly formed alveolar bone, periodontal ligament and cementum were regenerated without aberrant events. BMSC provided no added value to healing

Table 2 continued

Mericanin D <i>et al.</i> 2014 ⁽¹⁾	-PDLSC -P3 -undifferentiated 2×10^6	Gelfoam® +Gore-Tex®	SHEEP female (a, b) 7	-intrabony defects -5 mm in depth -no inflammation	(a) PDLSC+Gelfoam® (b) Gelfoam®	2 months	-Histological: HES, modified tetrachrome -Histomorphometric -Immunohistochemical: OPN, BSP, COL-I, αSMA, BrdU	PDLSC group shows a superior regeneration in bone and collagen fibers (similar to Sharpey's fibers) inserted in neocement
Mrozik KM <i>et al.</i> 2013 ⁽⁷⁾	-PDLC -P3 -undifferentiated 1×10^7	Gelfoam® +resorbable barrier membrane	SHEEP Merino ewes, 3–5 years old, $63.5\text{--}72.0\text{ kg}$ (a, b) 7	-dehiscence -10 mm in depth -no inflammation	(a) PDLC+Gelfoam® (b) Gelfoam® (c) empty	1 month	-Histological: HES, tetrachrome staining -Histomorphometric	Enhanced cementum regeneration following allogeneic PDLSC implantation
Nagahara T <i>et al.</i> 2015 ⁽⁸⁾	-BMSC -P3 -undifferentiated 2×10^7	βTCP/ atelocollagen	DOG Beagle, female, 10–14 kg (a, b) 20	-class III furcation defects -4 mm in depth -inflammation	(a) BMSC+β-TCP+atelocollagen (b) BMSC+atelocollagen (c) β-TCP+atelocollagen (d) atelocollagen	1 and 2 months	-Histological: HES, Azan and TRAP Staining -Immunohistochemical: OPN	The new cementum length in the groups (a, b) was higher at 4 and 8 weeks compared to control groups. And new connective tissue fibers were inserted into the cementum in both groups (a, b). The new bone volume was improved in group (a) but no in group (b)
Nakahara T <i>et al.</i> 2004 ⁽⁹⁾	-PDLSC -undifferentiated 3×10^5	type I (70–80%) and type III (20–30%) atelocollagen +ePTFE membranes	DOG Beagle, female, 10–12 kg (a, b) 6	-penetration defects -6×4 mm² -no inflammation	(a) PDLSC+atecollagen (b) empty	1 month	-Histological: HES, Masson's trichrome staining -Histomorphometric	Cement formation was a lot more significant with PDLSC. But no difference in bone formation between the groups
Nakajima R <i>et al.</i> 2014 ⁽²⁾	-derived stem cells -osteinductive medium	PGA+β-TCP	DOG Beagle (a, b) male, 1 year old, 10 kg	-intrabony defects -5×5 mm² -no inflammation	(a) DSC/PGA+β-TCP (b) PGA+β-TCP	2 months	-Histological: HES, Azan staining -Histomorphometric	A histologic analysis showed newly formed bone in both groups, whereas newly formed cementum-like tissue and Sharpey's fiber-like tissue were observed in the group (a) only.
Núñez J <i>et al.</i> 2012 ⁽⁶⁾	-CDC or PDLSC -P5-6 -undifferentiated ·CDC> 75×10^4 , PDLDC> 75×10^4	collagen	DOG Beagle, female, 50–56 months old 9–11 kg (a, b, c) 8	-intrabony defects -3×4 mm² -inflammation	(a) CDC+collagen (b) PDLSC+collagen (c) collagen	3 months	-Histological: Toluidine blue staining -Histomorphometric	Higher rates of cement found in test groups as well as a more significant gain in attachment and in conjunctive tissue. PDLSC or CDC failed to stimulate further the bone regeneration
Ozasa M <i>et al.</i> 2014 ⁽⁹⁾	-ASC -P4	fibrin gel	DOG Beagle, female, 50–56 months old 9–11 kg unknown	-class II furcation defects -4 mm in depth -inflammation	(a) ASC+fibrin gel® (b) fibrin gel	1, 5 months	-Histological: AZAN -Histomorphometric -Radiological: micro-CT	ASC transplanted sites have shown periodontal regeneration, including new alveolar bone, periodontal ligament and cementum formation with vertically inserted fibers

Table 2 continued

Paknejad M <i>et al.</i> 2015 ⁽⁷⁾	-BMSC -P3 -undifferentiated -2×10 ⁷ /default	Bio oss®	DOG Mongrel, male, 1-2 years old, 14-22 kg	9 (a, b)	-intrabony defects -4×4 mm ² -inflammation	(a) BMSC+Bio oss® (b) Bio oss®	2 months	-Histological: HES -Histomorphometric	Formation of new cementum and periodontal ligament were significantly higher in the test group. Whereas bone formation in the test and control groups were no statistically different
Simsæk SB <i>et al.</i> 2012 ⁽⁴⁾	-BMSC or Autologous cortical bone -undifferentiated -1×10 ⁷ /mL	PRP autologues	DOG Mongrel, 15 kg	6 (a, b, c, d, e)	-class II furcation defects -5×2 mm ² -no inflammation	(a) BMSC+PRP (b) autogenous cortical bone+PRP (c) PRP (d) autogenous cortical bone (e) no treatment	2 months	-Histological: HES -Histomorphometric	The regeneration of cement and alveolar bone was more significant in the groups (a, b, d), even if, in these 3 groups, BMSC showed the highest potential for periodontal regeneration.
Soroyama W <i>et al.</i> 2006 ⁽⁸⁾	-PDLC+SCAP -P1-3 -4×10 ⁶	Gelfoam®+HA+ β-TCP	PIG male, 4-8 months old, 20-40 kg	6 (a, b)	-alveolar socket -no inflammation	(a) [PDLC+SCAP/Gelfoam®]+HA +β-TCP (b) HA+β-TCP	1 month	-Histological: HES -Immunohistochemical: human mitochondria antibody -Radiological: CT	Periodontal regeneration observed in test groups: formation of a bio-root with a good compressive force
Suaid FF <i>et al.</i> 2011 ⁽⁸⁾	-PDLC -P2-3 -undifferentiated -3×10 ⁵	collagen sponge +absorbable membrane Goretex	DOG Beagle, 10-20 kg, 1.46±0.18 years old	7 (a, b)	-intrabony with class II furcation defects -5×2 mm ² -no inflammation	(a) PDLC+collagen (b) collagen	3 months	-Histological: HES -Histomorphometric	The results presented a better periodontal regeneration in the group with LAD cells
Suaid FF <i>et al.</i> 2012 ⁽⁸⁾	-PDLC -P2-3 -undifferentiated -3×10 ⁵	Collagen +2 absorbable membranes (GTR)	DOG Beagle, 10-20 kg, 1.46±0.18 years old	7 (a, b, c, d)	-intrabony with class II furcation defects -5×2 mm ² -no inflammation	(a) PDLC+collagen (b) collagen+GTR (c) GTR (d) empty	3 months	-Histological: HES -Histomorphometric	The group with LAD cells showed a better periodontal regeneration with a more significant bone and cement width
Tobita M <i>et al.</i> 2013 ⁽⁸⁾	-ASC -P2 -undifferentiated -1.5×10 ⁷ /mL	PRP gel	DOG Beagle, 9 or 10 months old, 8-10 kg	8 (a, b, c)	-intrabony with with class III furcation defects -5 mm in depth -no inflammation	(a) ASC+PRP gel (b) PRP gel (c) empty	2 months	-Histological: HES -Histomorphometric -Immunohistochemical: Osteocalcin -Radiological: X-ray	The results presented a better periodontal regeneration in test group. The group without treatment showed gingival invasion
Tsunanuma Y <i>et al.</i> 2011 ⁽³⁾	-PDLC or BMSC or Perosteal cells -P3 -osteoinductive medium -cell sheet	PGM/β-TCP/type I collagen	DOG Beagle, male, 10 kg	4 (a, b, c, d)	-intrabony defects -5×5 mm ² -no inflammation	(a) PDLC/PGM+ β-TCP/type I collagen (b) BMSC/PGM/β-TCP/type I collagen (c) Perosteal cells/PGM/β-TCP/ type I collagen (d) PGM/β-TCP/type I collagen	1 month	-Histological: HES, Azan staining -Histomorphometric -Immunohistochemical: neurofilament protein -Radiological: X-ray	PDL cells group shows a more significant cement thickness. Moreover, presence of correctly orientated collagen fibers, while they are slanted in the group (b) and parallel in the group (c). However, no difference in bone formation or in long junctional epithelium

Table 2 continued

Tsumanuma Y <i>et al.</i> 2016 ⁽³⁾	-PDLCSC -P5 -osteoinductive medium -cell sheet	PGA/β-TCP/ collagen +absorbable membrane (GTR) (TR)	DOG Beagle 8 (a, b, c)	-intrabony defects -5×5 mm ² -no inflammation	(a) PDLCSC autologous/PGA+ β-TCP+collagen (b) PDLCSC allogenic/PGA+ β-TCP+collagen (c) β-TCP+collagen	2 months	-Histological: Azan staining -Enzyme-linked immunosorbent assay: CRP, IL-10, IFN-γ, CD30 -Histomorphometric -Radiological: micro-CT	In the group (b), dense collagen fibers were observed, which attached perpendicularly to the cementum-like tissue. In the groups (a, c), collagen fibers were oriented obliquely or parallel to the root surface. There were no differences between the autogenous or allogenic groups in the histomorphometric analyses.
Wei N <i>et al.</i> 2010 ⁽⁰⁾	-BMSC -P3 -2×10 ⁷ /defect	Alginate hydrogel	DOG Beagle male, 6–10 months old, 5–10 kg	-class III furcation defects -5 mm in depth -no inflammation	(a) BMSC+hydrogel (b) empty	1, 5 months	-Histomorphometric: HES or Masson trichrome -Immunohistochemical: BrdU, α-SMA, osteocalcin	The transplanted BMSC migrated into the periodontal ligament, alveolar bone, cementum and blood vessels. BMSC transplantation has the potential to regenerate periodontal tissue.
Yang JR <i>et al.</i> 2013 ⁽⁷⁾	-Embryonic stem cells -P43-45 -GFP-label -1×10 ⁶ /defects	Collagen	MINIATURE PIG female 5 months old 25–30 kg	-classe II furcation defects -4×5–3 mm ³ -inflammation	(a) embryonic stem cells MSC+ collagen (b) collagen	3 months	-Clinical: PD, CAL -Histological: HES -Immunohistochemical: GFP, penistatin, aspirin, cementum attachment protein, osteopontin; osteocalcin, RUNX2	Embryonic cells to improve the regeneration of periodontal furcation defects is feasible
Yang KC <i>et al.</i> 2016 ⁽²⁾	-DPSC: P3-P5 -epithelial cells: P2-P31.76- 3.1×10 ⁶ -odontoblastic- induced DPSC: 20.3–33.2×10 ⁶ -osteoblastic- induced DPSC: 19.2–27.8×10 ⁶	gelatin chondroitin- hyaluronan	MINIATURE PIG 7 males, 10 females, 5–7 weeks old	-alveolar sockets -no inflammation	(a) cells layers+scaffold (b) scaffold (c) empty (d) without intervention	13.5 months	-Histological: HES -Immunohistochemical: DMP-1, OPN, COL-1, CK14, VEGF -Radiological: X-ray	This study demonstrated that a bioengineered cell/scaffold could achieve tooth regeneration. The regenerated tooth had crown, root, and pulp structures with enamel-like tissues, dentin, cementum, odontoblast-like cells, and periodontal tissues.
Zang S <i>et al.</i> 2016 ⁽³⁾	-BMSC -P3-5 -undifferentiated	-chitosan -anorganic bovine bone	DOG Beagles, 15 months old, 10–15 kg	-intrabony defects -4×7 mm ² -no inflammation	(a) BMSC+chitosan (b) BMSC+chitosan+ anorganic bovine bone (c) chitosan+anorganic bovine bone (d) chitosan (e) anorganic bovine bone (f) empty	2 months	-Histological: HES, Masson's trichrome staining -Histomorphometric -Immunohistochemical: osteocalcin -Radiological: micro-CT	BMSC associated with chitosan/anorganic bovine bone scaffolds could promote periodontal repair. The quantity of the newly formed bone and cementum in the groups (b, e) was significantly higher compared with the other groups.

Table 2 continued

Zhu B <i>et al.</i> 2017 ²⁵⁾	-PDlSC or BMSC (iliac or jaw-derived) -P2-4 -osteoinductive medium	treated dentine matrix (TDM)+ceramic bone (CA)	MINIATURE PIG 2 years old	10 (a) (b)	-intrabony defects -5.2x5 mm ² -no inflammation	(a) iBMSC+PDlSC+TDM+CA (b) iBMSC+PDlSC+TDM+CA	3 months	Both construct implantation, JBMSC or BMSC showed PDl-like tissue regeneration. However, the implantation of JBMSC seems to be more appropriate to form parallel collagen fibers and bone tissue
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HA, hydroxyapatite; TCP, tricalcium phosphate; PGA, polyglycolic acid; ePTFE, e-polytetrafluoroethylene; PRP, platelet-rich plasma; DPSC, dental pulp stem cells; PDlSC, periodontal ligament stem cells; DFSC, dental follicle stem cells; CDC, cementum derived cells; GMSC, gingival margin stem cells; SHED, stem cells from human exfoliated deciduous teeth; BMSC, bone marrow stem cells; ASC, adipose stem cells; MSC, mesenchymal stem cells; SCAP, stem cells from apical papilla; CAL, clinical attachment level; PI, plaque index; PD, probing depth; GR, gingival recession; BOP, bleeding on probing; HES, hematoxylin eosin stain; ECM, extracellular matrix; OPN, osteopontin; BSP, bone sialoprotein; Col, collagen; α SMA, alpha smooth muscle actin; GFP, green fluorescent protein; TRAP, tartrate-resistant acid phosphatase; CRP, serum c-reactive protein; IL-10, interleukin-10; INF- γ , interferon- γ ; CD, cluster of differentiation; VEGF, vascular endothelial growth factor; DMP-1, dentin matrix protein-1; CT, computed tomography; SEM, scanning electron microscopy.

Table 3 Comparative table of human studies in periodontal regeneration combining MSC and biomaterials

Reference	Cell type/ passage number/ differentiation/ number per defect	Biomaterial/H- membrane	Animal model/ species/sex/age/ weight	Number of defects per group	Defect type/size/ induced inflammation	Treatment groups	Observation period	Qualification of newly formed tissues	Results
Aimetti M <i>et al.</i> 2014 ¹¹⁾	-PDSC -P0 -undifferentiated	Collagen sponge (type III)	HUMAN male 56 years old	1	-intrabony defects -inflammation	DPSC+collagen sponge scaffold	6 and 12 months	-Clinical: CAL, PI, PD, GR -Radiological: dental X-ray	At the 12 months examination, the CAL gain amounted to 6 mm with a residual probing pocket depth of 3 mm. No apical displacement of the gingival margin was observed
Chen FM <i>et al.</i> 2016 ¹⁰⁾	-PDlSC -undifferentiated -cells sheets	Bio oss® +Bio-Gide®	HUMAN Male, female 26.05±4.44 30.04±7.90 years old	20 (a) 21 (b)	-intrabony defects ≥3mm in depth -inflammation	(a) PDlSC+Bio-oss® (b) Bio-oss®	3, 6 and 12 months	-Clinical: CAL, PD, GR -Radiological: dental X-ray	Significant difference between the experimental and the control groups was not observed
Dhote R <i>et al.</i> 2015 ⁴⁾	-P0 -rh-PDGF-BB stimulation -1x10% default	β-TCP	HUMAN 8 males and 6 females, 20-43 years old	12	-intrabony defects ≥5 mm in depth -inflammation	(a) allogenic human umbilical cord+β-TCP (b) open flap debridement	6 months	-Clinical: CAL, PI, PD, GR, BOP -Radiological: dental X-ray	Stimulated cells implantation resulted in a significant added benefit in terms of CAL gains, PI reductions greater radiographic defect fill and improvement in linear bone growth
Feng F <i>et al.</i> 2010 ³⁰⁾	-PDlC -P1 -undifferentiated	calcium carbonate	HUMAN male 25 and 42 years old	12 teeth Patient n°2 3 teeth Patient n°3 1 tooth	-intrabony defects ≥6 mm in depth -inflammation	PDlC+calcium carbonate	3-7/2 months	-Clinical: CAL, PD, GR, BOP, PI -Radiological: dental X-ray	PDlP implantation improve periodontal tissue regen, specifically marked by a significant decreased in gingival recession and increased attachment gain

Table 3 continued

Hernandez-Morajarz B <i>et al.</i> 2018 ⁽²⁾	-allogenic DPSC ·P3 ·undifferentiated ·5×10 ⁶ /defaut	Collagen sponge +teflon-coated titanium membrane	HUMAN male 61 years old	1	-intrabony defects -inflammation	Allogenic DPSC+collagen sponge	3, 6 months	-Clinical: PD, mobility -Radiological: Cone beam volumetric tomography; bone mineral density
Ki V <i>et al.</i> 2017 ⁽⁵⁾	-PDlSC ·P0 ·undifferentiated	Abgel® TM : gelatin sponge	HUMAN male 27 years old	1	-intrabony defects -9 mm in depth -inflammation	PDLSC+Abgel® TM	24 months	-Clinical: CAL, PI, PD, GR, relative attachment -Radiological: dental X-ray
Li Y <i>et al.</i> 2016 ⁽³⁾	-DPSC ·P3 ·undifferentiated	β-TCP	HUMAN female 30 and 38 years old	Patient n°1 1 tooth Patient n°2 1 tooth	-combined periodontal- endodontic lesions -5-6 mm in depth -inflammation	DPSC+β-TCP	1, 3, and 9 months	-Clinical: CAL, PI, PD, GR, BOP, furcation lesion, mobility -Radiological: dental X-ray
Okuda K <i>et al.</i> 2009 ⁽⁷⁾	-Perosteal cells ·P0 ·undifferentiated	HA+PRP	HUMAN female 53, 63 and 71 years old	3	-Intrabony defects -inflammation	Perosteal cells+HA+PRP	6 months	-Clinical: PD, CAL -Radiological: dental X-ray
Rosen PS <i>et al.</i> 2015 ⁽⁸⁾	-MSCs allografts (Osteocel, NuVasive) ·undifferentiated	deminerilized freeze-dried and freeze-dried bone allografts +BioXclude®	HUMAN Male and female 39-70 years old	6	-class III and IV furcation defects -inflammation	MSC+BioXclude®	6, 9, 24 or 30 months	-Clinical: furcation lesion -Radiological: dental X-ray
Yamada Y <i>et al.</i> 2005 ⁽¹⁾	-BMSC ·osteoinductive medium ·1×10 ⁷ /mL	PRP gel	HUMAN female, 54 years old	1	-intrabony defects -inflammation	BMSC+PRP gel	12 months	-Clinical: CAL, PD -Radiological: dental X-ray
Yamamoto K <i>et al.</i> 2008 ⁽⁶⁾	-periosteal cells ·P0 ·undifferentiated	HA+PRP	HUMAN 14 females and 1 male, 55.8-9.1 years old	15 (a, b)	-intrabony defects -6 mm in depth -inflammation	(a) periosteal cells+PRP+HA (b) HA+PRP	12 months	-Clinical: CAL, PD, GR, BOP, gingival inflammation -Radiological: dental X-ray

TCP, tricalcium phosphate; PRP, platelet-rich plasma; DFSC, dental pulp stem cells; PDLSC, periodontal ligament stem cells; BMSC, bone marrow stem cells; MSC, mesenchymal stem cells; CAL, clinical attachment level; PI, plaque index; PD, probing depth; GR, gingival recession; BOP, bleeding on probing.

The gingiva showed no signs of
inflammation, and depth of the periodontal
pocket and dental mobility both decreased.
Densitometry assays revealed an increase
in bone mineral density in the walls of
the defect

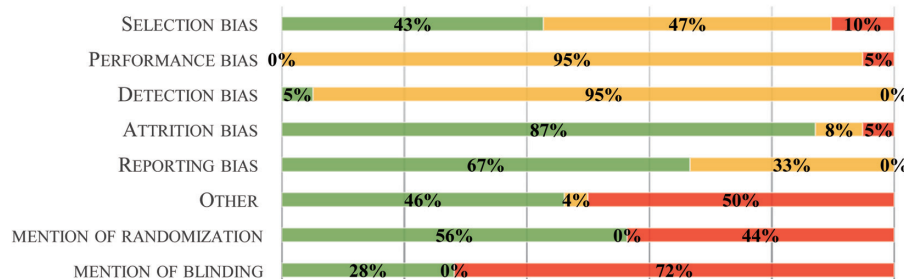
PDLSC implantation has resulted in
successful clinical and radiographic
parameters: CAL, decreased PD and
satisfactory defect fill of intrabony defects

most of the articles used intrabony defects (26 studies). It seemed relevant to underline that although periodontitis is an inflammatory disease, the defects models employed in preclinical studies were largely not inflammatory (in 26/39 studies, defects were surgically created). Then, the studies were categorized according to the type of MSC or biomaterial used. The periodontal ligament stem cells (PDLSC) were by far the most studied with 16 publications, whereas bone marrow stem cells (BMSC) were used in only 12 studies. As for scaffolds, collagen is used in most of the studies (10 studies). The general analysis of clinical studies has shown that studies mostly based on animals mainly used a combination of PDLSC and collagen.

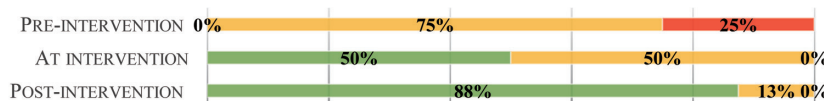
Risk of bias within studies

Figure 2 shows the overall results of the risk of bias assessment. Subgroup analyses was performed to assess the quality of preclinical studies (Fig. 2a), non-randomized human trials (Fig. 2b) and randomized human trials (Fig. 2c), using an adapted methodology. Firstly, the risk of bias assessment for preclinical animal studies was investigated. Establishing that only 44 and 72% of the studies do not mention randomization or blinding respectively, our data show a high score of unclear risk of bias for the selection, performance and detection items (47, 95 and 95%, respectively). Interestingly, the data outcomes were adequately addressed for 87% of the studies. Furthermore, the majority of the studies were free from selective outcome reporting (67%). Thereafter, non-randomized human trials were investigated for

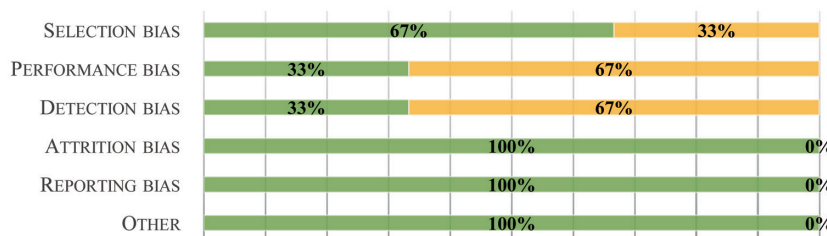
A. PRECLINICAL ANIMAL



B. NON-RANDOMIZED HUMAN TRIALS STUDIES



C. RANDOMIZED HUMAN TRIALS STUDIES



■ Low risk ■ Unclear risk ■ High risk

Fig. 2 Risk of bias assessment of included studies.

A, Risk of bias graph for animal studies, using the SYRCLE's tool, averaged per item.; B. Risk of bias graph for non-randomized human trials, using the ROBINS-I tool, averaged per item.; C. Risk of bias graph for randomized human trials, using the Cochrane Collaboration's tool, averaged per item. The green, yellow and red colors depict the percentages of studies with low, unclear or high risk of bias of the total number of assessed studies.

the quality of the methodology. Results revealed that 75 and 50% of the studies were marked for unclear risk of bias for pre-intervention and at-intervention items respectively. On the opposite, post-intervention items display a low risk of bias (88%). As expected for randomized human trials, our results did not show a high risk of bias. However, regarding selection bias, 33% of the included studies were marked for an unclear risk of bias. In addition, only 33% of the articles were considered to have a low risk of bias for its performance and detection. Finally, all studies were considered to have a low risk of bias for their attrition and reporting.

Synthesis of results

For each study, the relevant results are summarized for studies of preclinical animal models (Table 2) and for Human studies (Table 3). Overall analysis showed the association of MSC and biomaterial enhancement of periodontal regeneration for the majority of the studies.

DISCUSSION

The significant impact of periodontal disease on general health and the quality of life necessitates the need to regenerate the damaged tissue more effectively¹⁹. Consequently, the regeneration of bone, cementum and an effective periodontal ligament remains a major challenge. Periodontal regenerative medicine is considered as a promising treatment modality for future therapy. Along this line, the present systematic review intended to investigate the controversial results raised from the scientific literature on the efficacy of the periodontal regeneration strategies, using the association of MSC and biomaterials in pre-clinical animal models and humans.

Literature searches retrieved 50 studies. After a careful analysis, our results revealed that it was not possible to perform direct head-to-head comparisons of these studies as a result of variations between studies, in terms of the healing time after cell transplantation, the biomaterials applied, the defect type and size, the used cell types and passage number, and the number of cells per defect. Not surprisingly, no meta-analysis of the data could be carried out.

Indeed, the study of the risk of bias has revealed that poor reporting of animal studies in scientific publications is of serious concern. Preclinical animal studies and case reports are, in general, analyzed with less methodological rigour than trials. Key measures to avoid bias such as, randomization and blinding, were infrequently reported. This may lead to an overestimation of the effects of cells on periodontal regeneration compared to the group control system. This seriously hampers drawing reliable conclusions from animal studies. Despite these limitations, the combined analysis of the included studies still generated extra and valuable information that could not be derived from the individual analysis of studies.

Periodontal regenerative medicine is a multidisciplinary field combining biology and

engineering. In this context, the present discussion focuses first on MSC and second on biomaterials used for periodontal regeneration.

Stem cell biology has become an important field in regenerative medicine and tissue engineering therapy since the discovery and characterization of MSC. In particular, stem cells have great versatility at the level of tissue regeneration for many different characteristics and can modulate chronic inflammation, a central feature in periodontitis. Given the characteristics of these cells, they are considered a potentially useful tool for the efficient regeneration of periodontal tissues²⁰. Thus, one of the most important issues for clinical application of regenerative medicine approaches is the type of cell used.

Bone marrow has been the main source of MSC used for regenerative medicine. For several years now, BMSC has been the object of a lot of periodontal regeneration research, more often on animal models. To date, there exists only one case reported on a human suffering from chronic periodontitis. The authors concluded that the combination of BMSC and Platelet-rich Plasm (PRP) should entail a radiological and clinical improvement in terms of pocket depth, attachment gain, loss of bleeding on probing, and tooth mobility²¹. In addition, a randomized trial, focusing on the safety and the efficacy of regenerative treatment of infrabony defects using autologous BMSC, combined with collagen scaffolds enriched with fibrin glue is ongoing (NIH clinical trial registration number: NCT02449005). The combination of BMSC and PRP has also been a success on animal models in 3 different studies, each independently conducted by Simsek *et al.*, Hasegawa *et al.*, and Kawaguchi *et al.*²²⁻²⁴. Even though the periodontal regeneration was nearly complete in the study by Simsek *et al.*, both Hasegawa and Kawaguchi unfortunately concluded with incomplete tissue regeneration and particularly, that of the alveolar bone. With average positive results, the BMSC seems to enable an improvement in bone, periodontal ligament and cement regeneration^{25,26}. However, bone marrow suffers certain limitations that are related to its painful harvest and to the limited number of collectable cells. In light of these limitations, authors investigate other sources of adult tissues in order to collect stem cells. Within the orofacial area, several sources of MSC have attracted scientific interest, given their similarity to BMSC, their immunoregulatory capacity, and their minimally invasive harvest procedure²⁷.

Periodontal (PDL) derived cells

Thirty years ago, the concept that stem cells may reside in the periodontal tissues was put forward by Melcher²⁸. Not until 2004, PDLSC were first isolated and characterized as stem cells²⁹. Over the past several years, the number of animal studies on PDL-derived cells have been increasing. Not surprisingly, in the present systematic review, the majority, *i.e.* 40% of the 50 studies were dedicated to periodontal ligament-derived cells. Feng *et al.* were the first to transplant progenitor cells of the periodontal ligament combined

with calcium carbonate, into periodontal defects of 3 patients suffering from chronic periodontitis. The results concluded in a gain of clinical attachment with a decrease in pocket depth, a bone tissue regeneration and finally, a recession improvement over 72 months³⁰. Menicanin *et al.* have combined autologous PDLSC to a gelatin scaffold on sheep models. The results in the test groups were very encouraging due to a superior regeneration with the development of cement, bone and ligament structures³¹. In addition, Liu *et al.* and Ding *et al.* independently obtained the same types of results, with a near complete regeneration of the periodontal tissues after implanting a combination of autologous PDLSC to HA/βTCP in intrabony defects on porcine models over a period of 12 weeks^{32,33}. Moreover, in order to simplify the clinical protocol set for the dental surgeon in his everyday practice, some studies use periodontal ligament cells without isolating the stem cells beforehand. Combined with either a collagen scaffold, calcium carbonate, PGA, HA, β-TCP or hyaluronic acid, different authors agree that their cells shown a significant improvement in terms of the height and thickness of the bone, of cement and the periodontal ligament regeneration^{30,34-39}. A systematic review by Bright *et al.* on PDL-derived cells for periodontal regeneration reported that 12 out of the 17 included studies relate a statistically significant positive effect in periodontal regeneration¹⁴. A recent randomized clinical trial on 20 patients suffering from chronic periodontitis revealed that the implantation of PDLSC associated with Bio-Oss® over a period of 1 year significantly improved the alveolar bone height and the clinical parameters over time, but no significant differences between the implantation of PDLSC/Bio-Oss® and the implantation of only Bio-Oss® were found⁴⁰. In conclusion, even if the results are contradictory, the majority of the studies reveal a positive and promising effect on the regenerative potential of PDL-derived cells.

The use of dental pulp stem cells (DPSC) has shown some potential towards regenerating the periodontium. Interestingly, 3 independent case reports described an improvement of the clinical parameters of periodontitis by the implantation of DPSC with collagen scaffold^{41,42} or β-TCP⁴³. Although the results of several pre-clinical animal studies and case reports were promising, human randomized trials are required to evaluate the efficacy of those procedures in regenerating true periodontal defects.

The others stem cells used

The efficiency of cellular therapy using autologous gingival fibroblasts has been evaluated by Fawzy *et al.* when they were combined with gingival margin stem cells (GMSC) to Bio-Oss® or IL-1ra-releasing HA-ECM scaffold on a miniature pig subject. Whatever the biomaterial used, the team of researchers concluded with a reduction of pocket depth, a gain of clinical attachment and of bone density^{44,45}. In 2008, Yamamiya *et al.* conducted a study on 30 patients suffering from chronic periodontitis, where they combined periosteal cells with

PRP and HA over a period of 1 year⁴⁶. The results were positive from a clinical and a radiological point of view. Okuda *et al.* also studied the same combination in a dog model and reached the same conclusion⁴⁷. Adipose stem cells (ASC) has always seemed very appealing to researchers in periodontal regeneration. Tobita *et al.* and Ozasa *et al.* implanted ACS into periodontal defects on canine models with PRP⁴⁸ or fibrin gel⁴⁹, respectively. The morphometric, histologic, immuno-histologic and radiological analysis confirmed a superior bone, cement and ligament formation in the cell group.

Interestingly, a previous systematic review and meta-analysis, which included a large number of studies using only animal models (sheep, dog, minipigs, rats, mice), provided evidence for the enhancement of periodontal regeneration by the implantation of either PDL-derived cells or BMSC^{50,51}.

To have a chance of achieving periodontal regeneration, the cells need to be delivered and stabilized on the defect by a biomaterial. The concept of a scaffold is based on a biomimetic strategy, capable of incorporating and releasing molecules, and permitting cell to cell and cell to matrix interactions. The choice of the releasing method will depend on the type of cell population released and on the type of defect; specifically, the number of alveolar bone walls involved. When defects are larger, outcomes may improve when cells are associated with bone substitute or cell sheets attached with PGA are transplanted onto the tooth root surface, after which the bone defects are filled with bone substitute^{36,38,39,52,53}. On the opposite, gel scaffold might be used when lesions are retentive. As shown in this systematic review, clinical studies mainly use a scaffold made of collagen. Collagen materials may be particularly useful due to their biocompatibility, resorbability, cell occlusiveness and their capability of promoting wound healing. Collagen offer a safe scaffold material because of occurring naturally and being involved in numerous physiological processes. Although collagen can be constituted into various forms such as fibers, sheets, hydrogels, and sponges¹¹. Most biomaterials have some drawbacks such as follows: unpredictable cell-biomaterial interactions, non-homogeneous biodegradation, immune reaction, and low efficiency cell seeding. Injectable scaffolds, easily applied without invasive surgery, can improve cell retention, distribution and more importantly activate *in situ* cell proliferation and differentiation. Hydrogels are not only biocompatible with a high resorption level but they show a better cell retention than other injectable biomaterials, as confirmed by a number of studies⁵⁴. They can be prepared from alginate, chitosan, collagen or cellulose, seeded with cells and then gelation can be initiated by changing the temperature, pH, cross-linking or radical polymerization. The goal is to obtain a high cell retention followed by a good integration capacity and a high level of surviving cells with reduced side effects and minimal stress for the patient¹².

The present systematic review achieves, for the first time, a comprehensive analysis of the scientific literature concerning the periodontal regenerative

medicine. To date, current data indicate that MSC associated with suitable scaffolds may provide beneficial effects on periodontal regeneration in preclinical animal models and humans. In particular interest, the human studies suggest that there are no adverse effects of such interventions^{40,42}.

Although clinical trials are promising, future animal studies are still needed to determine the suitable association of MSC and biomaterial for periodontal regeneration and to characterize the type of cementum, and the organization of the periodontal ligament fiber that is regenerated. Furthermore, these studies should also compare periodontal regenerative medicine with the gold standard therapies used to repair periodontium.

CONCLUSION

Several approaches using MSC for regenerating damaged periodontium are under study with varying degrees of clinical applications. Given the heterogeneity of the studies concerning the periodontal regenerative medicine, narrative reviews are insufficient. A systematic approach appears essential to provide guidance to support future studies and should provide information that can be generalized. Our results indicate that MSC may provide beneficial effects on periodontal regeneration. The present systematic review supports crucial information for the implementation of regenerative medicine strategies in clinical practice in the future.

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CONFLICTS OF INTEREST

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