Multipotent adult progenitor cells on an allograft scaffold facilitate the bone repair process

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Abstract
Multipotent adult progenitor cells are a recently described population of stem cells derived from the bone marrow stroma. Research has demonstrated the potential of multipotent adult progenitor cells for treating ischemic injury and cardiovascular repair; however, understanding of multipotent adult progenitor cells in orthopedic applications remains limited. In this study, we evaluate the osteogenic and angiogenic capacity of multipotent adult progenitor cells, both in vitro and loaded onto demineralized bone matrix in vivo, with comparison to mesenchymal stem cells, as the current standard. When compared to mesenchymal stem cells, multipotent adult progenitor cells exhibited a more robust angiogenic protein release profile in vitro and developed more extensive vasculature within 2 weeks in vivo. The establishment of this vascular network is critical to the ossification process, as it allows nutrient exchange and provides an influx of osteoprogenitor cells to the wound site. In vitro assays confirmed the multipotency of multipotent adult progenitor cells along mesodermal lineages and demonstrated the enhanced expression of alkaline phosphatase and production of calcium-containing mineral deposits by multipotent adult progenitor cells, necessary precursors for osteogenesis. In combination with a demineralized bone matrix scaffold, multipotent adult progenitor cells demonstrated enhanced revascularization and new bone formation in vivo in an orthotopic defect model when compared to mesenchymal stem cells on demineralized bone matrix or demineralized bone matrix–only control groups. The potent combination of angiogenic and osteogenic properties provided by multipotent adult progenitor cells appears to create a synergistic amplification of the bone healing process. Our results indicate that multipotent adult progenitor cells have the potential to better promote tissue regeneration and healing and to be a functional cell source for use in orthopedic applications.

Keywords
Multipotent adult progenitor cell, demineralized bone matrix, tissue engineering, angiogenesis, osteogenesis, bone

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Introduction
Regenerative medicine emphasizes the use of stem cells in conjunction with biologic and synthetic scaffolds. The application of stem cells to a wound site can substantially improve the time, quality, and overall extent of healing.1–6 It is widely known that production and maintenance of bone tissue are mediated largely by a cascade of molecular signals that are released by and acted upon lineage specific stem cells. These cells can either differentiate or participate in further signal conduction with growth factors and hormones to facilitate bone remodeling.7 Resident bone cells can provide signals to osteoprogenitor cells through pathways including, but not limited to, Wnt, transforming growth factor-beta (TGF-β), and bone morphogenic protein (BMP).7–12

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Osteogenesis is traditionally considered the most important element for new bone formation; however, it is widely known that angiogenesis plays an important role in bone repair as well. New blood vessel formation allows for the migration of cells and necessary nutrients to the site of injury. Studies have shown that promotion of vessel formation in an injury model can influence bone healing and that health factors that negatively impact neovascularization (i.e. smoking or diabetes) can cause delayed fracture healing or result in nonunions. Furthermore, the literature has demonstrated that vascular endothelial growth factor (VEGF) is crucial to the bone remodeling process. Studies have shown that both VEGF and BMPs increase the differentiation of pre-osteoblasts and that they elicit a synergistic effect on bone formation.

Many orthopedic injuries involve major trauma to the skeletal system and require surgical intervention. Therapeutic approaches to treating damaged or diseased tissue can include a combination of hardware, bioactive large molecules (i.e. growth factors and cytokines), viable cells, and/or natural or synthetic scaffolds. While hardware and scaffolds assist in maintaining structural support and load-bearing integrity, bioactive large molecules and viable cells provide signals for the host to upregulate new tissue formation and stimulate the healing process. One of the most common cell sources for orthopedic applications is bone marrow–derived mesenchymal stem cells (MSC). MSC are operationally defined as plastic adherent fibroblastic-like cells capable of differentiating along mesodermal lineages, including bone.

While MSC are established, and their applications are well published in the orthopedic field, there may be alternative cell types possessing greater therapeutic implications. One such population, termed multipotent adult progenitor cells (MAPC), was first described in 2002 by Jiang et al. MAPC are nonhematopoietic stem cells derived from the bone marrow stroma. MAPC have a broader lineage differentiation capacity than MSC, generating cells of the mesenchymal lineage, as well as endothelium, hematopoietic cells, hepatocyte-like cells, and neuroectoderm-like cells. In addition to their broader differentiation potential, MAPC can proliferate without obvious signs of senescence and can be expanded to over 70 passage doublings while remaining cytogenetically normal.

To date, research has focused predominantly on the immunomodulatory properties of MAPC as well as their therapeutic efficacy in models of myocardial infarction and hypoxic ischemia. MAPC have been shown to differentiate along mesodermal lineages, including undergoing osteogenic differentiation in vitro; however, MAPC have not been fully investigated for their use in orthopedic applications. A previous study performed in a heterotopic model demonstrated increased osteoinductivity of a MAPC-loaded demineralized bone matrix (DBM) scaffold. While the authors presented the ability of MAPC to undergo and/or promote ossification in vivo, the study did not utilize an injury model in an orthotopic site. Another limitation of the study was the absence of an injury-induced inflammatory response, which is significant in the bone healing process. Studies have shown that stem cells may require an inflammatory stimulus to initiate the healing response. While an initial inflammatory response is normal and may trigger advantageous cellular responses, prolonged or persistent inflammation can negatively impact the healing process. MAPC possess demonstrated immunomodulatory properties and the ability to attenuate a local host immune response upon implantation. Lehman et al. established that endothelial cells exhibit a reduced production of vascular cell adhesion molecule (VCAM), E-selectin, and intercellular adhesion molecule (ICAM) when co-cultured with MAPC; the attenuated presence of these proteins reduces neutrophil binding to endothelial cells. This phenomenon leads to decreased endothelial activation that may reduce inflammation and neutrophil infiltration.

This study further investigates the role of MAPC in the orthopedic milieu by applying them in an orthotopic, segmental defect model, while comparing their performance to the current MSC standard. In this study, MAPC isolated from human bone marrow demonstrated a select angiogenic protein release profile which surpassed that of MSC in vitro. In vivo, MAPC seeded onto DBM and implanted into a fibular defect contributed to wound closure and promoted enhanced vascularization. In conjunction with improved neovascularization, treatment groups with MAPC demonstrated increased bone healing. These results suggest a synergistic relationship between angiogenic and osteogenic elements that may accelerate the bone healing process.

Materials and methods

Cell isolation and culture

MAPC used in this study were isolated as previously described by Yasuhara et al. and cultured as described by Boozer et al. MAPC were characterized according to the methods of Sohni and Verfaillie. MAPC were cultured and expanded at 37°C and 3% O2. Passage numbers ranging from 2 to 4 were used for all experiments. MAPC isolated from 10 donors were validated for consistency in morphology, growth rates, surface markers, and cytokine...
expression (data not shown). Bone marrow–derived MSC (Normal, Human, ATCC®PCS-500-012) were purchased from ATCC (Manassas, VA) and cultured according to manufacturer instructions. MSC used in this study were selected as a model system and have been verified by ATCC to expand to 15 population doublings while still maintaining characteristics of primary MSC. These characteristics include morphology, growth curves, differentiation potential, surface marker expression, immunosuppression, and tube formation.59 These data are comparable to findings published on isolated primary bone marrow–derived MSC.60–63

Osteogenic differentiation

One million MAPC were seeded onto fibronectin (5 ng/mL)-coated flasks and cultured in MAPC maintenance medium (Lonza, Basel, Switzerland) for 3 days at 37°C and 3% O₂. After 3 days, the medium was replaced by osteogenic differentiation medium.64 Osteogenic differentiation medium consisted of high-glucose Dulbecco’s modified Eagle’s medium (DMEM; Life Technologies, Carlsbad, CA), dexamethasone (Sigma–Aldrich, St Louis, MO), ascorbic acid 2-phosphate (Sigma–Aldrich), β-glycerophosphate (Sigma–Aldrich), fetal bovine serum (Life Technologies), and penicillin–streptomycin (Life Technologies). Cells were cultured at 37°C and 21% O₂ for an additional 14, 21, and 28 days, with medium changes every 3–4 days.

Adipogenic differentiation

MAPC were prepared as described in the “Osteogenic differentiation” section with the substitutions of adipogenic differentiation medium65 and Oil Red O staining (Sigma–Aldrich). Adipogenic differentiation medium contained α-minimum essential medium (MEM; Life Technologies), fetal bovine serum, hydrocortisone (Sigma–Aldrich), isobutylmethylxanthine (Sigma–Aldrich), indomethacin (Sigma–Aldrich), penicillin–streptomycin, and dexamethasone.

Chondrogenic differentiation

One million MAPC were seeded onto fibronectin (5 ng/mL)-coated flasks and cultured in maintenance medium for 3 days at 37°C and 3% O₂. After 3 days, cells were transferred onto a round bottom 96-well tissue culture plate at a density of 4.5 × 10⁵ cells/well and cultured in chondrogenic differentiation medium64 (high-glucose DMEM, dexamethasone, ascorbic acid 2-phosphate, proline (Sigma–Aldrich), sodium pyruvate (Sigma–Aldrich), penicillin–streptomycin, Insulin transferrin selenium (ITS) + premix (Corning, Corning, NY), and TGF-β1 (Sigma–Aldrich)). MAPC were aggregated by centrifugation at 500g for 5 min and then returned to the hypoxic incubator. After 24h, the medium was changed and aggregates were gently released from the sides and bottom of the wells by pipetting. Aggregates were cultured for 14, 21, and 28 days, with media changes every 3–4 days. At the end of the culture period, the aggregates were fixed in formalin, dehydrated, embedded in paraffin, and stained with toluidine blue (Sigma–Aldrich).

Qualitative alkaline phosphatase staining

MAPC or MSC were cultured in control medium or osteogenic medium for 8 days. Alkaline phosphatase (ALP) staining was performed using a naphthol AS-MX phosphate and fast red violet B salt-based kit (Sigma–Aldrich). Briefly, cells were fixed in citrate-buffered acetone and rinsed with water. Cells were exposed to the alkaline dye mixture for 30 min, rinsed with water, visualized using light microscopy, and imaged. Images were taken at 10× magnification.

Quantitative ALP staining

MAPC or MSC were cultured in osteogenic medium for 8 days. Supernatant media were collected and analyzed for the presence of ALP using QuantiChrom™ ALP Assay Kit (BioAssay Systems, Hayward, CA). Briefly, media were exposed to the supplied working solution, and the optical density was read at 405 nm at t=0 min and t=4 min.

Qualitative alizarin red staining

MAPC or MSC were cultured in control or osteogenic medium for 21 days. Alizarin red staining was performed according to standard protocol.66 Briefly, cells were rinsed with phosphate-buffered saline (PBS) and fixed in 10% formalin. Cells were stained with alizarin red (Sigma–Aldrich) for 20 min, rinsed with water, and visualized using light microscopy. Images were taken at 10× magnification.

Quantitative calcium assay

MAPC or MSC were cultured in osteogenic medium for 21 days. Calcium production was analyzed using a calcium reagent set (Pointe Scientific, Canton, MI). Briefly, cells were lysed with 0.5 N HCl to expose the mineral to the acid, and the volume was collected. The samples were incubated with calcium reagents for 10 minutes at room temperature, and the absorbance was read at 570 nm.

Angiogenic protein analysis

MAPC or MSC were plated at a density of 1 × 10⁴ cells/well in a 24-well tissue culture plate. After 24h, the medium was removed and replaced with fresh serum-free
medium. Following an additional 24 h at 37°C, 3% O₂ (MAPC), or 21% O₂ (MSC), the media were harvested for use in an enzyme-linked immunosorbent assay (ELISA). IL-8, CXCL-5, VEGF, and GRO-α ELISAs (R&D Systems, Minneapolis, MN) were performed according to the manufacturers’ instructions and normalized to total protein levels using the BCA Protein Assay Kit (Thermo Fisher Scientific, Waltham, MA).

**Matrigel tube formation assay**

Human umbilical vein endothelial cells (HUVEC; Lifeline Technology, Frederick, MD) were cultured in standard medium until they reached 70%–80% confluence. Matrigel (Becton Dickinson, Franklin Lakes, NJ) was added to the wells of a µ-angiogenesis slide (ibidi, Verona, WI) and allowed to polymerize for 30 min at 37°C. A total of 1 × 10⁴ HUVEC were added to each well in 25 µL of medium, along with 25 µL of positive control medium (basal maintenance medium supplied by the manufacturer with growth factors for vessel formation), negative control medium (media devoid of growth factors), or conditioned medium from MAPC or MSC. Wells were imaged at 2, 4, and 6 h.

**Surgical methodology**

Male athymic rats (7–8 weeks old) purchased from Harlan Laboratories (Indianapolis, IN) were used for this study. The animals were acclimated for 48 h prior to surgery. On the day of surgery, animals were anesthetized and surgically prepared. Surgical procedure and postoperative care were conducted according to the established protocol approved by the Institutional Animal Care and Use Committee of the University of Florida. Briefly, the fibula was accessed through a lateral skin incision and blunt dissection of the musculature. A 4-mm segmental defect in the fibula was created unilaterally. Defects in experimental groups were treated with a DBM scaffold seeded with MAPC (MAPC + DBM scaffold) or MSC (MSC + DBM scaffold) at a concentration of 175,000 cells/cm³ of DBM, while defects in the control group received DBM scaffold only. All DBM scaffolds originated from the same donor lot to ensure consistency. Musculature was sutured, and skin wounds were closed using skin clips. Animals were sacrificed at 14 and 28 days post-implantation for histological evaluation of new vessel and bone formation.

**Histological methodology**

After the explants and surrounding tissue were removed, the specimens were fixed, decalcified, and embedded in paraffin using standard protocols. Specimens from the 14-day time point were sectioned transversely, while those from the 28-day time point were sectioned longitudinally to better expose the defect areas for evaluation. Four sections representing different depths into the defect were collected and stained with hematoxylin and eosin (H&E). Blood vessel quantification was performed on the 14-day specimens, and bone repair was evaluated on the 14- and 28-day specimens.

**Histological evaluation**

All histological evaluation, analysis, and reporting were performed by an outside contract laboratory. The magnitude of vascular network formation and bone healing at the fracture site was evaluated using a semiquantitative severity scoring system with a 0- to 5-point scale width, in which higher scores correlated with normal tissue architecture or improved healing. Tables 1–3 illustrate the comprehensive scoring criteria.

**Immunohistochemistry**

Immunohistochemical detection of von Willebrand factor (vWF) protein was performed on sections of formalin
fixed, paraffin-embedded rat fibulas. Sections were deparaffinized in xylene and then rehydrated through a graded alcohol series. Antigen retrieval was performed for 10 min at 37°C using a Proteinase K solution (Life Technologies). Nonspecific binding was blocked with 10% goat serum (Life Technologies) and 1% bovine serum albumin (BSA; Sigma–Aldrich) in tris-buffered saline and Tween 20 (TBS) for 1 h at room temperature. Sections were then incubated overnight with anti-vWF antibody (Abcam, Cambridge, England) at a dilution of 1:200 in the blocking buffer. Alexa Fluor 488 goat anti-rabbit IgG (ThermoFisher Scientific, Waltham, MA) was added for 1 h at room temperature, protected from exposure to light. Slides were counterstained with 4′,6-diamidino-2-phenylindole (DAPI; Life Technologies), and coverslips were mounted with Permount Mounting Medium (Thermo Fisher Scientific). Protein-antibody complexes were detected using an Olympus DSU-IX81 spinning disc confocal microscope (Olympus, Shinjuku, Tokyo, Japan). A total of 6–10 images were taken per stack, and images were deconvolved using the nearest-neighbor method. Maximum intensity z-projection was used to obtain two-dimensional (2D) images. Signal intensity and exposure duration were consistent across all groups.

**Statistics**

All quantitative assays were performed in at least triplicate, and the mean value was used. Data are presented as mean ± standard deviation (SD) with \( p < 0.05 \) indicative of significance, unless otherwise specified. If normality passed, a Student’s \( t \) test (for comparison of two groups) or a standard analysis of variance (ANOVA; for comparison of three or more groups) was used, followed by Tukey’s post hoc test. If normality failed, a nonparametric Kruskal–Wallis ANOVA on ranks was used followed by a Mann–Whitney \( U \) post hoc test. Data were analyzed using SigmaStat software (Systat Software, Inc., Chicago, IL).

**Results**

**MAPC differentiate along the mesoderm lineages**

To complement surface marker characterization data as previously described,③ MAPC were cultured in specific medium that promotes differentiation along osteogenic, chondrogenic, and adipogenic lineages. Osteogenic potential was demonstrated by visualization of calcium deposition in the extracellular matrix using alizarin red (Figure 1(b)). Adipogenesis was confirmed by cytoplasmic lipid droplets and intracellular lipid vesicle formation via Oil Red O staining (Figure 1(c)). Finally, chondrogenesis was evidenced by positive staining for glycosaminoglycans using toluidine blue staining (Figure 1(d)).

Figure 1. Multipotent differentiation potential of MAPC: (a) MAPC were cultured in control medium and (b–d) selected differentiation media for 21 days: (b) differentiated MAPC displaying calcium deposition with alizarin red staining, (c) differentiated MAPC exhibiting positive staining for lipids using Oil Red O stain, and (d) differentiated MAPC with positive staining for proteoglycans with toluidine blue. Images a–c are at 20× magnification. Image d is at 10× magnification.

Biphasic organization, indicative of a chondrogenic phenotype, was observed as well. Positive staining was not observed in any of the control conditions (Figure 1(a)). MSC differentiation was performed by ATCC.⑤

**MAPC qualitatively express more ALP when compared to MSC**

MAPC and MSC were cultured in control and osteogenic media for 8 days. Qualitative staining indicated low levels of ALP expression (dark purple staining) by MAPC in control medium, which became abundant when the cells were exposed to osteogenic conditions (Figure 2(a) and (b)). MSC cultured in control medium did not express ALP (Figure 2(c)), with only mild expression following culture in osteogenic medium, mostly restricted to sparse and random areas (Figure 2(d)).

**MAPC quantitatively express more ALP when compared to MSC**

MAPC and MSC were cultured in control media for 8 days. The quantitative assay indicated significantly higher expression of ALP by MAPC when compared to MSC (Figure 2(e)).

③ MAPC differentiate along the mesoderm lineages

⑤ MAPC qualitatively express more ALP when compared to MSC

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MAPC produce mineral deposits

No evidence of mineral deposition was observed in MAPC, and MSC cultured under control conditions for 21 days (Figure 3(a) and (c)). However, calcium-containing mineral deposits were present in the extracellular matrix of both cell types under osteogenic conditions after 21 days (Figure 3(b) and (d)); both cell cultures exhibited similar levels of expression qualitatively; however, quantitative data demonstrated a significant increase in expression in the MAPC group (Figure 3(e)).

MAPC exhibit increased angiogenic protein expression compared to MSC

An angiogenic cytokine array was performed to qualitatively evaluate the secretion of angiogenic signaling molecules in MAPC and MSC (data not shown). We subsequently quantitatively measured the expression of four commonly known angiogenic markers: VEGF, GRO-α, IL-8, and CXCL-5, using ELISAs. MAPC produced IL-8 (Figure 4(a)) and GRO-α (Figure 4(b)) at significantly higher levels than MSC, while CXCL-5 (Figure 4(d)) was highly produced by MAPC, but undetectable for MSC. VEGF expression was comparable between MAPC and MSC (Figure 4(c)). A matrigel tube formation assay was performed to determine the impact of these significant differences in protein secretion on endothelial cells. The endothelial cells treated with MAPC conditioned medium exhibited a dense tube-like formation within 6h, while those in the MSC conditioned medium group had only very sporadic tube formation at the same time point (Figure 4(e)).
MAPC induce new blood vessel formation in a fibular defect after 14 days

MAPC + DBM scaffold or MSC + DBM scaffold were implanted into a bone void, and the resultant blood vessel formation was examined after 14 days (Figure 5). Histological evaluation revealed that MAPC + DBM scaffold groups developed significantly more blood vessels when compared to the DBM scaffold control and MSC + DBM scaffold groups (Figure 5(a)–(c) and (g)). To highlight these differences in blood vessel formation, slides were stained with vWF (Figure 5(d)–(f)). An increased number of blood vessels were evident in defects treated with MAPC + DBM scaffold (Figure 5(b) and (e)) compared to those treated with MSC + DBM scaffold and the DBM scaffold alone (Figure 5(a), (c), (d), and (f)). In addition, vessels in the MAPC + DBM scaffold group were larger in size and exhibited more mature morphology (Figure 5(b) and (e)).

MAPC + DBM scaffold promotes increased bone repair after 14 and 28 days

The extent of repair in an acute long bone defect was evaluated using an osteogenic approach with MAPC + DBM scaffold and MSC + DBM scaffold as treatment groups. Tables 1 and 2 illustrate the scoring criteria for woven and lamellar bone. After 14 days, the MAPC + DBM scaffold treatment groups had significantly more woven and lamellar bone (Figure 6(d) and (e), respectively) when compared to the scaffold-only group. In addition, the average total bone (Figure 6(f))

Figure 4. In vitro angiogenic potential of MAPC and MSC. (a–d) ELISA analysis using conditioned media from MAPC and MSC in control conditions; (a) ELISA analysis of interleukin 8 (IL-8) secretion in MAPC versus MSC, (b) GRO-α secretion in MAPC versus MSC, and (c) VEGF secretion in MAPC versus MSC. (d) CXCL-5 secretion in MAPC versus MSC. CXCL-5 was below detectable limits of the assay for MSC (n = 3 for each, *p < 0.01). (e) Matrigel tube formation assay with human umbilical vein endothelial cells (HUVEC) exposed to MAPC and MSC conditioned media, n = 3, 6-h time point.
was increased in the MAPC + DBM scaffold group when compared to either the DBM scaffold-only or the MSC + DBM scaffold group.

Furthermore, at 28 days, mineralized callus containing developing areas of bone marrow was detected in the MAPC + DBM scaffold group with minimal fibrous tissue (Figure 7(b)). The MAPC + DBM scaffold treatment group demonstrated significantly higher deposition of woven bone, when compared to the MSC + DBM scaffold treatment group (Figure 7(d)). While the MSC + DBM scaffold group also contained woven bone, it was to a lesser degree than in the MAPC + DBM scaffold group, with substantial fibrous tissue between proximal and distal ends of the fibular defect in the MSC + DBM scaffold group (Figure 7(a)–(c)).

Additionally, higher levels of marrow organization (Table 3) are present in the MAPC + DBM scaffold
treatment group (Figure 7(e)), further supporting the accelerated bone healing demonstrated in the 14-day data.

**Discussion**

Currently, the stem cell landscape has expanded from uncharacterized stromal cells to a variety of identifiable cell types (embryonic cell, MSC, marrow-isolated adult multilineage inducible (MIAMI) cell, very small embryonic-like (VSEL) cell, etc.) intended for a multitude of applications including cardiovascular, neural, and musculoskeletal repair.\(^{67-70}\) One of the more recently studied stem cells, MAPC, has the common ability to differentiate along the osteogenic lineage in vitro.\(^{25,27,58}\) We aimed to confirm the potential role of MAPC in an orthopedic setting by expanding upon recent published data. To this end, we selected an in vivo bone defect model that would evaluate osteogenesis, as well as angiogenesis. The rat fibular defect model was chosen for its associated low levels of inflammation, which can provide an environment conducive to cell signaling.\(^{50}\) Previous studies have demonstrated that 14- and 28-day time points provide sufficient time to evaluate revascularization and bone healing, respectively, in this model.\(^{50,67}\)

The osteogenic potential of MAPC was verified through an evaluation of osteogenic markers, specifically ALP and degree of mineralization. ALP, a well-known early marker of the osteogenic phenotype, was measured in vitro. MAPC mineral deposition was demonstrated after 21 days in culture via staining with alizarin red. Both ALP and mineral deposition were expressed at higher levels in MAPC when compared to MSC. In vitro angiogenic results demonstrated that MAPC-secreted proteins promoted neovascularization through their effect on HUVEC in a tube formation assay, to a significant level over MSC. Specific angiogenic proteins, such as GRO, IL-8, and CXCL-5, which may contribute to the increased tube formation in vitro and the upregulated neovasculature observed in vivo,\(^{41-44,71}\) were all expressed at significantly higher levels in MAPC when compared to MSC.
To determine whether in vitro angiogenic and osteogenic potential translated to in vivo efficacy, we performed a 28-day study in a rat fibular defect model with MAPC seeded onto DBM scaffolds (which provided the necessary osteoconductive and osteoinductive elements for bone formation). This constitutes the first time that MAPC have been used in a clinically relevant orthopedic application. In addition, this study compared the overall healing potential of MAPC to that of the more commonly used MSC. The DBM scaffold control allowed us to isolate the contribution of the cellular component to osteogenesis and angiogenesis. Healing was first indicated by the presence of increased vasculature in the MAPC + DBM scaffold treatment group at 14 days in comparison to the scaffold-only or MSC + DBM scaffold controls. This was evaluated quantitatively and confirmed via fluorescent-conjugated antibody staining for vWF. It has previously been demonstrated that MAPC possess angiogenic properties that make them ideal for use in cardiovascular applications. Although it is rarely emphasized, angiogenesis is an important factor in the bone healing cascade, allowing for delivery of cells and nutrients to the damaged tissue during the healing process. Lack of nutrient transport between damaged tissue and the healthy surrounding tissue can often compound the disruptive effects of a bone injury. Deficient vasculature and subsequent impeded revascularization may slow down the healing process and can lead to partial or incomplete healing such as nonunions. Before osteogenesis can occur, vessel healing and revascularization must begin, making this an essential element in bone repair.

Once in vivo neovascularization was confirmed, repair and bridging in a bone defect model were evaluated. New bone formation after 14 days provided evidence of an osteogenic response at the defect site and defects with implanted MAPC + DBM scaffold demonstrated increased bone repair when compared to MSC + DBM scaffold and scaffold-only controls. We speculate that the increase in angiogenic factors resulted in enhanced neovascularization in lieu of an angio-inductive element, allowing for improved nutrient availability as well as an influx of osteoprogenitor cells. These factors likely contributed to the increased bone healing present at the 14-day time point in the MAPC + DBM scaffold group.

Based on the results of this study, it is likely that MAPC + DBM scaffold groups are advancing more rapidly through the stages of bone healing when compared to the MSC + DBM scaffold and scaffold-only groups. At 14 days, there was increased neovascularization, as well as
a higher degree of vessel maturity (Figure 5) demonstrated. This is likely a result of the elevated levels of angiogenic factors secreted by MAPC, which subsequently increase the development of blood vessels within the defect. This increase in vasculature resulted in an increase in total bone healing in the MAPC + DBM scaffold group at 14 days (Figure 6). Between 14 and 28 days, the MAPC + DBM scaffold group underwent a significant increase in woven bone, while the MSC + DBM scaffold group maintained a low level, and the DBM scaffold-only group experienced a minor increase (Figure 7). This is a possible indication that MAPC + DBM scaffold groups are progressing more rapidly through the callus ossification stage, in which the vessels that matured in the first 2 weeks provided an influx of osteoprogenitor cells, which then mineralized the cartilage callus and formed woven bone. Additionally, there is early indication of a transition from woven to lamellar bone (Figure 6) in the MAPC + DBM scaffold group, which may suggest that this group is advancing into the bone remodeling stage, the final stage of the ossification process. This is further supported by the early stages of marrow development at 28 days (Figure 7). Figure 8 represents a speculative mechanism of healing of MAPC on DBM scaffold.77

Figure 8. Proposed Mechanism of MAPC+DBM Scaffold bone healing. Data in this manuscript indicate high degrees of neovascularization and vessel maturity by 2 weeks. This increase in vasculature accelerates formation of the cartilage callus, which is then mineralized to form woven bone within the 2 week period. After 2 weeks, the increase in woven bone begins to plateau, and by 4 weeks there is increased lamellar bone and early marrow development. These results suggest that during the first 2 weeks of healing, defects in the DBM+MAPC scaffold treatment group progress through the callus formation and callus ossification stages, and at 4 weeks the woven bone is being resorbed and replaced with primary lamellar bone, which is characteristic of an advance into the bone remodeling stage. Bone illustrations were adapted from Frohlich M (77).
result in the more rapid total repair of the defect. The results of this study demonstrate MAPC + DBM scaffold as a promising therapeutic for clinical use in orthopedic applications. Future studies will focus on understanding the signaling mechanisms that drive the strong angiogenic and osteogenic effects of MAPC, the potential early callus formation and chondrogenic effects, cell and scaffold interactions, and the pathways involved.

Conclusion
An in vivo model paired with multiple in vitro assays demonstrated the potential contribution of MAPC at various stages of bone healing. This is the first study to show that MAPC + DBM scaffold exhibit osteogenic and angiogenic properties in an orthotopic, fibular defect model. Furthermore, specific angiogenic factors that may impact revascularization during bone regeneration have been identified, both in vitro and in vivo. Finally, the osteogenic and angiogenic properties of MAPC were compared to those of the more commonly studied MSC. We have demonstrated that MAPC are a promising therapeutic for clinical use in orthopedic applications and have set the foundation for future studies to investigate the mechanisms by which MAPC contribute to the bone healing cascade. This study has implications for the development of synthetic, natural, and composite scaffolds that may benefit from an osteogenic cellular component for the treatment of damaged bone tissue.

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