Three-Dimensional Engineered Bone from Bone Marrow Stromal Cells and Their Autogenous Extracellular Matrix

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Most bone tissue-engineering research uses porous three-dimensional (3D) scaffolds for cell seeding. In this work, scaffold-less 3D bone-like tissues were engineered from rat bone marrow stromal cells (BMSCs) and their autogenous extracellular matrix (ECM). The BMSCs were cultured on a 2D substrate in medium that induced osteogenic differentiation. After reaching confluence and producing a sufficient amount of their own ECM, the cells contracted their tissue monolayer around two constraint points, forming scaffold-less cylindrical engineered bone-like constructs (EBCs). The EBCs exhibited alizarin red staining for mineralization and alkaline phosphatase activity and contained type I collagen. The EBCs developed a periosteum characterized by fibroblasts and unmineralized collagen on the periphery of the construct. Tensile tests revealed that the EBCs in culture had a tangent modulus of $7.5 \pm 0.5$ MPa at 7 days post-3D construct formation and $29 \pm 9$ MPa at 6 weeks after construct formation. Implantation of the EBCs into rats 7 days after construct formation resulted in further bone development and vascularization. Tissue explants collected at 4 weeks contained all three cell types found in native bone: osteoblasts, osteocytes, and osteoclasts. The resulting engineered tissues are the first 3D bone tissues developed without the use of exogenous scaffolding.

Introduction

Approximately one million surgeries are performed each year in the United States that require bone grafts to replace tissue damaged by disease or extensive trauma. Several limitations are associated with grafting, such as graft availability, donor site morbidity, and immune rejection. Because of these complications, strategies are being developed to engineer bone tissue in vitro for bone replacement.

Current approaches to engineer bone involve the design of a three-dimensional (3D) scaffold that promotes the differentiation and proliferation of osteogenic cells and the deposition and mineralization of an osteogenic extracellular matrix (ECM). The scaffold design rubrics also include the ability to withstand physiological loads in vivo and eventual incorporation into the native tissue or degradation during the course of tissue development. Polymers such as poly(lactic-co-glycolic acid), poly(propylene fumarates), and poly(caprolactones) provide a matrix that promotes cell adhesion and migration, allow for the deposition and mineralization of osteogenic ECM in vitro, and have predictable degradation rates but lack the mechanical properties needed to withstand the loads placed on natural bone in vivo. Hydroxyapatite and β-tricalcium phosphates are ceramics used for bone scaffolding that also promote cell adhesion and proliferation and when implanted have shown positive results with regard to bone regeneration in vivo. However, the brittle nature of ceramics inhibits their use in healing large defects. Polymer–ceramic composite scaffolds such as calcium phosphate salts embedded in poly(caprolactones) have been designed to mitigate the problems with using each material alone, but a significant percentage of cells fails to attach to the composite scaffold because of limited surface-to-volume ratio. Single-layer cell sheets grown from bone marrow stromal cells (BMSCs) and wrapped around composite scaffolds have recently been shown to form constructs that resemble bone in vitro and in vivo. However, this method still involves the use of an exogenous scaffolding that must incorporate into native tissue.

BMSCs are multipotent cells that can differentiate in vitro in response to chemical signals and generate and mineralize their own autogenous ECM. BMSCs can be easily isolated from autologous sources and therefore serve as an attractive candidate for tissue engineering. Because a specific bone

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marker does not exist, an engineered tissue is characterized as being bone according to a number of criteria. The presence of alkaline phosphatase (ALP), an enzyme that cleaves phosphate ions from organic molecules, is a precursor to mineralization of bone and thus an early sign of bone formation. Bone is composed predominantly of type I collagen mineralized with hydroxyapatite crystals and contains osteocalcin, so the presence of these molecules further substantiates differentiation of BMSCs toward bone. The absence of type II collagen, the predominant protein of cartilage, is often used to exclude differentiation to cartilage. The structural properties of bone, including cells with an osteogenic morphology and nucleation and growth of mineralized structures through matrix vesicles or intrafibrillar mineralization, can be monitored using light and electron microscopy. The periosteum of bone is fibrous tissue found on the bone surface that contains fibroblasts, osteoprogenitor cells, and unmineralized type I collagen. Osteoprogenitor cells differentiate into osteoblasts during bone growth and remodeling. The periosteum can be identified through microscopy because of the elongated shape of fibroblasts and osteoprogenitor cells, versus the round presentation of osteoblasts. The presence of rows of osteoblasts between the fibrous periosteum and the mineralized bone core is an indication of a functioning periosteum because the osteoprogenitor cells are actively differentiating to osteoblasts.

In this study, we developed a method to engineer 3D bone-like tissue structures solely from BMSCs and their autogenous ECM using an approach similar to one we previously developed for tissue engineering of soft tissues and their inter-

**Methods**

**BMSC isolation and expansion**

All animal care and animal surgeries were in accordance with The Guide for Care and Use of Laboratory Animals (Public Health Service, 1996, National Institutes of Health Publication No. 85-23); the University of Michigan’s Committee for the Use and Care of Animals approved by the experimental protocol. Under aseptic conditions, bone marrow was collected from the femur and tibia of female Fisher 344 rats. Briefly, the soft tissues of the leg were removed from the femur and tibia, both ends of the bones were detached, and the marrow flushed out using a syringe with a 25-gage needle filled with Dulbecco’s modified Eagle medium (DMEM; Gibco, Rockville, MD). The marrow was vortexed and then centrifuged at 480 g for 5 min using a ThermoForma General Purpose Centrifuge (Waltham, MA). The pellet was resuspended in 10 mL of growth medium (M1) consisting of DMEM with 20% fetal bovine serum (Gibco), 6 ng/mL basic fibroblast growth factor (bFGF; Peprotech, Rocky Hill, NJ), 10^{-8} M dexamethasone (dex; Sigma-Aldrich, St. Louis, MO), and 1% antibiotic-antimycotic (Gibco), and plated into 100-mm-diameter tissue culture dishes. The dishes were kept in an incubator at 37°C, 95% humidity, and 5% carbon dioxide. After 48 h, the non-adherent cells were removed by replacing the M1. The adherent BMSCs were cultured to 80% confluence, at which time cells were enzymatically removed from the plate using a 0.25% trypsin-ethylenediaminetetraacetic acid solution (Gibco) and passed. Cells were plated onto construct dishes during the third to fifth passages.

**Construct dish preparation**

Tissue culture plastic dishes, 35 mm in diameter (BD Biosciences, San Jose, CA), were filled with 1.5 mL SYLGARD (type 184 silicone elastomer; Dow Chemical, Midland, MI). The polymer was allowed to cure for 3 weeks before use. The SYLGARD was coated with 3 mg/cm² natural mouse laminin (Invitrogen, Carlsbad, CA) by filling dishes with 3 mL of a 9.6 mg/mL laminin solution in Dulbecco’s phosphate buffered saline (DPBS; Gibco). The laminin solution was evaporated overnight in a biosafety cabinet. Dishes were rinsed once with DPBS and then filled with 1 mL of DMEM containing 20% fetal bovine serum and 1% antibiotic-antimycotic. The dishes were then sterilized via exposure to ultraviolet radiation (wavelength, 253.7 nm; bulb G30T8) in a biological safety cabinet for 60 min, then kept in an incubator for 5 to 8 days. After the incubation, the medium was aspirated from the dish, and BMSCs were seeded onto each dish in 2 mL of M1 supplemented with 0.13 mg/mL L-ascorbic acid-2-phosphate (asc-2-phos; Sigma-Aldrich) and 0.05 mg/mL L-proline (Sigma-Aldrich). There were 30 dishes in total, with an initial cell density of 200,000 BMSCs/dish. The cells were fed M1 supplemented with asc-2-phos and proline every 2 days until confluence was reached. Once the cells reached confluence, two minutiens pins, 0.2 mm in diameter and 1 cm long (Fine Science Tools, San Francisco, CA), were pinned onto the cell monolayer 1.5 cm apart. At confluence, the M1 was switched to a second medium (M2), consisting of DMEM with 7% horse serum (Gibco), 0.13 mg/mL of asc-2-phos, 0.05 mg/mL of L-proline, and 2 ng/mL of transforming growth factor beta (TGF-β; Peprotech). Ten of the 30 plates also contained 6 mg/mL of bFGF and 10^{-8} M dex and received the TGF-β at confluence (Bone Group 1), another 10 plates were also supplied the bFGF, dex, and TGF-β, but the TGF-β was administered 5 days after confluence (Bone Group 2). The remaining 10 plates received TGF-β at confluence but did not receive bFGF or dex (Bone Group 3). Table 1 lists the medium formulations used in this study. There were no differences in the ability of the cells in each population to form bone-like constructs, and the resulting constructs had similar histological structure and mechanical properties; therefore, results were combined from the three sets. Seven days after construct formation, the diameter of the samples was measured using a reticle in the eyepiece of a Nikon SMZ1500 dissection microscope (Melville, NY).

**Histochemical and immunofluorescence staining**

Seven days after 3D construct formation, the engineered bone-like constructs (EBCs) were mounted on a holder using tissue freezing medium (Triangle Biomedical Sciences,
Durham, NC) and immersed in −80°C isopentane. The frozen samples were sliced longitudinally to a thickness of 9 μm to 12 μm using a Microm HM 500 cryostat system (Heidelberg, Germany). The slides were then used for histological staining for light microscopy or immunofluorescent staining. For histochemical staining, tissue sections were fixed with methanol and stained for calcification with Alizarin Red or hematoxylin and eosin to observe tissue structure. The remaining sections were fixed in acetone and stained for ALP activity.

Immunofluorescent staining was performed to detect the presence of collagen I, collagen II, and osteocalcin. Frozen sections were fixed with methanol for 5 min and rinsed three times with DPBS. Sections were then blocked for 30 min with Ham’s F-12 containing 1% donkey serum. The concentrations of each of the antibodies were as follows: 5 μg/mL of rabbit anti-rat collagen I (Abcam Inc., Cambridge, MA), 5 μg/mL of mouse anti-riat collagen II (Calbiochem, Darmstadt, Germany), and 10 μg/mL of mouse anti-rat osteocalcin (Abcam). Samples were then rinsed three times with Ham’s F-12 and blocked again in Ham’s F-12 containing 5% donkey serum at 37°C for 10 min. The secondary antibodies (5 μg/mL) were then applied to the sections for 1 h as follows: Alexa Fluor 488 donkey anti-rabbit immunoglobulin (IgG) (Molecular Probes, Eugene, OR) for collagen I, Alexa Fluor 555 donkey anti-mouse IgG (Molecular Probes) for collagen II, and Alexa Fluor 488 donkey anti-mouse IgG (Molecular Probes) for osteocalcin. Positive and negative control stainings were performed using wheat germ agglutinin lectin and primary antibody omission, respectively (not shown). Nuclei were stained with Prolong Gold antifade reagent with 4’,6-diamidino-2-phenylindole (DAPI; Invitrogen). A Nikon Eclipse TS100 microscope equipped with an X-Cite 120 Fluorescence Illumination System was used to image the histochemically and immunofluorescently marked sections. Images were captured using a Diagnostic Instruments Spot Insight Color camera (Sterling Heights, MI).

Transmission electron microscopy

Samples were fixed for TEM in 2.5% glutaraldehyde (Electron Microscopy Sciences, Hatfield, PA) in 0.1 M of Sorenson’s buffer, pH 7.4, for 24 h at 4°C. Constructs were then thoroughly rinsed with Sorenson’s buffer and post-fixed with 1% osmium tetroxide in Sorenson’s buffer for 2 h. Samples were then rinsed with double distilled water and stained with 8% uranyl acetate in double distilled water for 1 h. Constructs were dehydrated in a graded series of ethanol, treated with propylene oxide, and embedded in Epon 812. Longitudinal ultra-thin sections, 70 nm thick, were prepared and stained with uranyl acetate and lead citrate. The sections were examined using a Philips CM100 electron microscope (Bothell, WA) at 60 kV. Images were digitally recorded using a Hamamatsu ORCA-HR digital camera system (Hamamatsu City, Japan) operated using AMT software (Advanced Microscopy Techniques Corp., Danvers, MA).

Mechanical properties

Tensile tests were performed on EBCs engineered in vitro 7 days (n = 4) or 6 weeks (n = 3) after 3D construct formation. Tests were performed using a custom tabletop tensiometer designed in our laboratory, attached to a Nikon SMZ1500 dissection microscope. The sample was immersed in a bath of DPBS, and the ends were grasped using clamps. The dimensions of the sample were measured before each test for cross-sectional area calculation using a reticle in the eyepiece of the microscope. Beads 25 μm in diameter (Interactive Medical Technologies, Irvine, CA) were brushed on the surface of the sample as position markers for digital image correlation analysis of tensile strain. Samples were stretched at a strain rate of 0.01/s. Force was measured using a custom force transducer and monitored using LABVIEW software (National Instruments, Austin, TX); the resolution of the force transducer used for testing the EBCs was 1 mN. Images of the sample were captured using a Basler camera (Exton, PA) attached to the Nikon dissection microscope at a frequency of 2.5/s during the test to follow the location of the digital image correlation markers. The bead position from each image was recorded using LABVIEW software (Melville, NY) with a resolution of 5 μm, and length between beads was calculated by subtracting the position of one bead from another. Nominal stress was calculated using the following equation:

\[ \sigma = \frac{F}{A_0} \]
Where $\sigma$ is nominal stress, $F$ is force, and $A_0$ is the initial cross-sectional area of the sample. Nominal strain was calculated using the following equation:

$$
\varepsilon = \frac{l - l_0}{l_0}
$$

Where $\varepsilon$ is the nominal strain, $l$ is the current length between beads, and $l_0$ is the original length. Moduli were obtained by calculating the slope of the tangent of stress versus strain plots at the maximum strain before failure.

**Implantation**

Seven days after 3D construct formation, EBCs were placed into a piece of sterile TYGON silicone tubing (United States Plastics Corp., Lima, OH) that had an inner diameter of 1.6 mm, an outer diameter of 3.2 mm, and a length of 1.5 cm. Fisher 344 rats were anesthetized using 0.001 mL of Nembutal sodium solution (pentobarbital sodium injection; Ovation Pharmaceutical Inc., Deerfield, IL) per gram of the animal’s weight. The silicone tubes containing the constructs were then implanted between the biceps femoris and the quadriceps of the left leg of the host animals. The silicone tubing was used to identify the engineered tissue during explantation.

Four weeks after implantation, constructs were removed from the animal. The diameter of the explants was measured using electronic calipers. The samples were then frozen and mounted in tissue freezing medium and stored at $-80^\circ$C or fixed in 2.5% glutaraldehyde and mounted in paraffin for histological studies.

**X-ray diffraction**

Wide-angle X-ray scattering diffraction was performed on EBC ($n = 2$) samples 6 weeks in culture post-3D construct formation. The bone constructs were removed from their medium and allowed to dry. The engineered tissues were placed on a single-crystal silicon substrate and examined using a Bruker (D8 Discover) diffractometer (Madison, WI) using Cu Kα radiation and a source voltage of 40 kV. The samples were investigated over a 2θ range of 23° to 55° with 20 counts recorded per degree. Scattering resulting from the substrate was subtracted, and resulting peaks were identified using the hydroxyapatite Powder Diffraction File from the Joint Committee on Powder Diffraction Standards (Swarthmore, PA).
Results

BMSCs were cultured on construct dishes in a medium that induced osteogenic differentiation (M1). Approximately 2 to 4 days after plating, the cells reached confluence and were switched to a second medium (M2) containing a lower serum concentration and TGF-β, and two minutien pins were attached to the culture dish as constraint points. After a sufficient amount of ECM was produced, the tissue monolayer lifted from the substrate and contracted radially. The placement of the two constraint points inhibited full contraction, and the monolayer formed into a cylindrical tissue construct of approximately 1.5 cm in length and an elliptical cross-section with major and minor axes of 1.0 ± 0.07 mm and 0.78 ± 0.13 mm, respectively (Fig. 1a). EBC formation was complete within 14 days of plating the BMSCs onto the cell culture dishes. The resulting engineered tissue was completely solid, which suggests that the cells within the EBCs remodeled the tissue during and after construct formation. The EBCs were kept in culture 7 days or 6 weeks after formation of 3D structure before being fixed for histological and structural characterization.

At 7 days post-3D EBC formation, constructs were sectioned longitudinally for histological investigation. Hematoxylin and eosin (H&E) staining revealed that the self-assembled EBCs were composed of dense and fibrous regions (Fig. 1b). The bulk of the tissue stained positive for ALP activity (Fig. 1c), and calcium deposits were located throughout the samples, as seen using Alizarin Red staining (Fig. 1d). The EBCs stained intensely throughout for type I collagen (Fig. 1e) and lacked type II collagen (Fig. 1f), which is consistent with the constitution of native bone. At this point, osteocalcin could not be detected within the EBCs (Fig. 1g). The DAPI-stained cells were fairly uniformly dense throughout the section (Fig. 1h). TEM verified that the EBCs contained osteoblasts (Fig. 2a) in a collagenous matrix undergoing mineralization. The diameter of the collagen was 29 ± 4 nm, which corresponds to the diameter of type I collagen found in the femur. Mineralization was noted to occur through matrix vesicles and intrafibrillar calcification (Fig. 2b, c). Elongated fibroblast-like cells were found on the periphery of the EBCs within axially aligned unmineralized type I collagen (Fig. 2d, e). This structure could be interpreted as the onset of periosteum development. Wide-angle X-ray scattering diffraction peak locations for the EBCs corresponded with those of hydroxyapatite standard (ICDD: 01-073-0293), verifying that the crystals seen in the TEM micrographs were indeed hydroxyapatite (Fig. 3).

The EBCs were able to maintain their size and shape after pins were removed from the edges. When the constructs were lifted and held horizontally at one end with forceps, they did not visibly deflect under their own weight or deform under the pinching loads of the forceps. Unlike (unmineralized) soft tissue constructs that we have engineered in our laboratory that contract longitudinally upon constraint pin removal and cannot support an axial compressive load, the EBCs resist compressive and bending deformation. Tension tests performed on EBCs in culture at 7 days and 6 weeks post-3D EBC formation revealed maximum tangent moduli of 7.5 ± 0.5 MPa (n = 4) and 29 ± 9 MPa (n = 3), respectively. The cross-sectional area remained the same between 7 days and 6 weeks post-3D construct formation. The tangent moduli are normalized with respect to the cross-sectional area of the EBCs; therefore, the increase in stiffness between the two time points is a measure of EBC phenotype development and not of physical growth.

At 7 days post-3D construct formation, the EBCs were implanted for 4 weeks between the biceps femoris and quadriceps of Fisher 344 rats. While implanted, the EBCs grew and remodeled so that the resulting explant was cylindrical, with a circular cross-section. The diameter of the explant was 1.6 ± 0.3 mm, equaling the inner diameter of the...
silicone tubing they had been placed in during implantation (Fig. 4a). H&E staining of the demineralized explants revealed a structure that appeared similar to that of native bone (Fig. 4b, c). Osteocytes in lacunae and blood vessels were seen throughout the construct (Fig. 4b, c). Further development of a periosteum-like structure was seen after implantation, as indicated by the fibrous tissue along the edge of the bone explants and the neighboring osteoblast-like cells (Fig. 4b). Alizarin Red staining of explant sections showed greater amounts of calcification (Fig. 4d) than the construct engineered in vitro (Fig. 1D). The explanted constructs contained type I collagen and lacked type II (Fig. 4e, f). After implantation, osteocalcin was present within the bone explants (Fig. 4g). DAPI staining (Fig. 4h) showed higher cell densities at regions of lower calcification (Fig. 4d, g vs h). This phenomenon is similar to that in native tissues, in which the cell density is lower in mineralized bone than in surrounding fibrous tissues. TEM of the demineralized explants showed a periosteum-like structure containing osteoprogenitor cells and rows of osteoblast-like cells (Fig 5a). Osteocytes in lacunae were seen within the bone matrix (Fig 5b). Osteoclasts were found throughout the mineralized tissue region (Fig 5c, d). The osteoclasts were identified via their multinucleation (Fig 5c), ruffled borders, and a plethora of vacuoles used by the cell for storage of resorbed material (Fig. 5d).

Discussion

We have developed a method to engineer 3D bone-like constructs from only BMSCs and their autogenous ECM. After 7 days of post-3D construct formation, the EBCs exhibited ALP activity and contained mineralized type I collagen. The mechanical properties of the EBCs improved over time in vitro; tangent stiffness increased by a factor of four over a 5-week period. No significant physical size change occurred in vitro from 7 days to 6 weeks in culture after 3D construct formation, indicating phenotype development due to tissue remodeling or increased collagen or mineral production. The phenotype of the EBCs continued to advance during implantation in vivo. The explants contained a vascularized bone-like structure with osteoblasts, osteocytes, and osteoclasts. The cell density was in mineralized regions than in the surrounding fibrous areas. The explants contained osteocalcin, which was not detected before implantation, and qualitatively stained more intensely for mineralization. The explants also had a functional periosteum-like tissue containing osteoprogenitor cells that were undergoing differentiation to osteoblasts.

The requirements for bone formation in vitro from BMSCs are ascorbic acid, dex, and an organic phosphate. These three components were administered to the BMSCs during the cell proliferation stage of the EBC culture (Table 1). Ascorbic acid maintains connective tissue and regulates ATPase, ALP, and protein synthesis in cultures of osteoblasts. Dex, a synthetic glucocorticoid, stimulates osteoblastic and adipogenic differentiation from BMSCs. Phosphates provide phosphate ions for matrix mineralization. In a study published by Maniatisopoulos et al., BMSCs were plated on tissue culture plastic to monitor bone formation from BMSCs in two dimensions. The BMSCs were cultured in an osteogenic medium containing ascorbic acid, dex, and an organic phosphate to form bone nodules. Similar to our EBCs, these nodules exhibited ALP activity, were composed of mineralized type I collagen and osteocalcin, and contained osteoblasts and osteocytes. In this previous study, control dishes that lacked dex contained 5 times more cells than dishes that received dex. It is hypothesized that the addition of bFGF to the culture medium allowed for monolayer for-
bFGF is commonly used as a potent mitogen for many types of mesenchymal cells. In bone cultures, this cytokine increases mineralization, ALP activity, and the concentrations of bone-specific markers such as calcium and osteocalcin when administered to BMSCs in an osteogenic medium. The mitogenic effects of bFGF in addition to dex in the present study may have increased proliferation, thus allowing for monolayer formation rather than the formation of nodules. Control dishes that received M2 lacking TGF-β formed cell monolayers that lifted from the substrates but did not contract into cylindrical constructs (data not shown); thus, TGF-β is the factor that controls 2D versus 3D construct formation. Although the overall effects of this growth factor on BMSCs are not fully known, it is generally used in culture to stimulate collagen production and matrix maturation and to induce chondrogenic differentiation from BMSCs. In bone cultures, TGF-β regulates osteoblast replication and migration, increases ALP activity, and stimulates collagen production and matrix maturation in bone cultures derived from osteogenic cells and BMSCs. We are performing studies to elucidate the effects of TGF-β on 3D bone construct formation with the hypothesis that TGF-β increases the rate of collagen production at an early stage of EBC development and before full osteogenic differentiation.

After 4 weeks in vivo, the EBCs grew to equal the size of the tubing they was placed into before implantation. It is unknown whether the osteoblasts and osteocytes seen in the EBC explants were derived from the cells present in the EBCs in vitro or from precursors from the native blood supply. If some of the osteoblasts and osteocytes developed from the native tissue, this is an indication that the construct provides an optimal scaffold for bone cell infiltration and differentiation even though it was placed in an ectopic site. Explants contained a significant amount of osteocalcin, but this protein was not found in the EBCs before implantation. Osteoclasts were also found within the mineralized matrix of the EBC explants. The observation of osteoprogenitor cells actively differentiating into osteoblasts and the presence of osteoclasts demonstrates that the EBCs actively grew and remodeled in vivo in a manner similar to that of native bone.

FIG. 4. Characterization of engineered bone-like construct (EBC) explants. (a) Photograph of explant after 4 weeks of implantation, scale bar = 5 mm. (b) Hematoxylin and eosin (H&E) staining of edge of sample, osteoblasts (arrow) differentiating from osteoprogenitor cells of FP and invading engineered bone (EB). (c) H&E of center of demineralized engineered bone shows osteocytes (black arrows) and blood vessels (white arrows). Scale bar of (b) and (c) = 62.5 μm. (d) Alizarin red staining for calcification, (e) collagen I, (f) collagen II, (g) osteocalcin, (h) and 4',6-diamidino-2-phenylindole staining for nuclei. Scale bar (d) through (h) = 250 μm. Color images available online at www.liebertonline.com/ten.
Before implantation, the periphery of the EBCd contained fibroblast-like cells and unmineralized collagen. This phenomenon is unique to this study and has not yet been reported in 2D bone nodules or in bones engineered with exogenous scaffolding. This structure could be interpreted as the onset of periosteum development. After implantation for 4 weeks, the engineered tissue developed a functional periosteum, as indicated by the presence of osteoprogenitor cells, and rows of osteoblasts (Fig. 5A). If the structure on the periphery of the EBC before implantation was the onset of the periosteum that continued to develop when implanted in vivo, it would indicate fibrogenic and osteogenic differentiation from a single cell source in vitro within the same culture environment. This could suggest that the BMSCs were differentiating because of the cues delivered through the medium and preferentially as a result of their location relative to other cells in an engineered construct in addition to chemical signals. Mechanically constraining the contractile monolayer induces tensile strain along the major axis of the constructs and preferentially orients the collagen fibers in this direction. These are the first 3D, scaffold-less tissues developed from bone marrow stem cells that demonstrate phenotype advancement in vitro and in vivo.

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