In vitro Apatite Mineralization, Degradability, Cytocompatibility and in vivo New Bone Formation and Vascularization of Bioactive Scaffold of Polybutylene Succinate/Magnesium Phosphate/Wheat Protein Ternary Composite

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Purpose: A bioactive and degradable scaffold of ternary composite with good biocompatibility and osteogenesis was developed for bone tissue repair.

Materials and Methods: Polybutylene succinate (PS:50 wt%), magnesium phosphate (MP:40 wt%) and wheat protein (WP:10 wt%) composite (PMWC) scaffold was fabricated, and the biological performances of PMWC were evaluated both in vitro and vivo in this study.

Results: PMWC scaffold possessed not only interconnected macropores (400 μm to 600 μm) but also micropores (10 μm ~ 20 μm) on the walls of macropores. Incorporation of MP into composite improved the apatite mineralization (bioactivity) of PMWC scaffold in simulated body fluid (SBF), and addition of WP into composite further enhanced the degradability of PMWC in PBS compared with the scaffold of PS (50 wt%)/MP (50 wt%) composite (PMC) and PS alone. In addition, the PMWC scaffold containing MP and WP significantly promoted the proliferation and differentiation of mouse pre-osteoblastic cell line (MC3T3-E1) cells. Moreover, the images from synchrotron radiation microcomputed tomography (SRmCT) and histological sections of the in vivo implantation suggested that the PMWC scaffold containing MP and WP prominently improved the new bone formation and ingrowth compared with PMC and PS. Furthermore, the immunohistochemical analysis further confirmed that the PMWC scaffold obviously promoted osteogenesis and vascularization in vivo compared with PMC and PS.

Conclusion: This study demonstrated that the biocompatible PMWC scaffold with improved bioactivity and degradability significantly promoted the osteogenesis and vascularization in vivo, which would have a great potential to be applied for bone tissue repair.

Keywords: polybutylene succinate, composite scaffold, cytocompatibility, osteogenesis, vascularization

Introduction

Bone tissue regeneration and reconstruction involve the use of biomedical materials to repair damaged/diseased bone tissue to restore biological functions for patients who suffer from osteoporosis, deformities, bone fractures, tumors, etc.1,2 Implantable biomedical materials with excellent biological performances and osteogenic bioactivity exhibit great potential for bone repair applications.3,4 As
a biomedical material for bone regeneration, bioactivity and biodegradability have crucial influences on cell responses as well as osteogenic bioactivity. Therefore, research and development of biodegradable biomedical materials with good bioactivity for bone regeneration have been paid more attention for many years.

Polybutylene succinate (PS), an aliphatic thermoplastic polyester, possesses a range of desirable properties, such as good biocompatibility, degradability, flexibility and processability. Notably, the degradation products of PS are demonstrated to be nontoxic and able to enter the metabolic cycles of bioorganisms. Despite these advantages, the low bioactivity, slow degradation rate, not synchronizing the new bone formation of a biomaterial still does not meet the requirement of bone tissue regeneration. Consequently, improving the bioactivity and degradability are needed for the further wide application of PS for bone repair.

Studies have revealed that magnesium ion (Mg\(^{2+}\)) plays critical roles in bone remodeling and skeletal tissue development of the human body. Growing evidence shows that Mg\(^{2+}\) may improve bone mineral density and affect bone fragility, and lack of Mg\(^{2+}\) can, definitely, affect all stages of skeletal metabolism, leading to slower bone growth and osteoporosis. In the past few years, Mg-containing bioactive materials have received increasing attention for bone tissue repair applications. Several Mg-based biomaterials for bone repair were reported, such as Mg-associated alloys, Mg-containing bioactive glasses and their coatings, and Mg-substituted calcium phosphate biocermics/biocerements. Unfortunately, the Mg-related inorganic scaffolds, such as magnesium phosphate (MP), even though having amazing similarity in bioactivity to commonly commercial available calcium phosphate scaffolds, generally shows a slow degradation rate, slightly unsynchronizing the formation of new bone tissue.

Natural-origin polymers, such as collagen, chitosan, and implant protein, are the most important raw materials for the development of biomedical materials. Wheat protein (WP) as an interesting natural polymer with excellent properties including good biocompatibility, hydrophilicity, and great biodegradability has been applied in many areas, such as coating, encapsulation, and other medical applications. In a previous study, it was reported that the attachment and proliferation of osteoblasts on WP were better than poly(lactic acid) (PLA). In addition, WP has been formulated with PLA and electrospun for biomedical applications.

Substantially, natural bone tissue consists of inorganic material (apatite) and organic compounds (collagen). Thus, to mimic the natural bone, the biocomposites containing bioactive inorganic materials and organic polymers have been widely researched and developed, and have advantages over individual inorganic materials and polymer. Based on the advantages of PS, MP, and WP, in this study a novel scaffold of ternary composite (PMWC) containing 50 wt% PS, 40 wt% MP and 10 wt % WP was prepared. The objective of this study is to develop a bioactive and degradable scaffold of ternary composite with good biological performances for bone regeneration and reconstruction. Therefore, the in vitro bioactivity, degradability, and cytocompatibility and the in vivo osteogenesis and vascularization properties of the PMWC scaffold were investigated.

**Materials and Methods**

**Fabrication and Characterization of Composite Scaffolds**

MP was synthesized by using a chemical precipitation method. Briefly, magnesium nitrate (MN) and ammonium dihydrogen phosphate (ADP) were dissolved in deionized water, respectively. The MN solution was added dropwise to the ADP solution while stirring to produce white precipitates, meanwhile, pH value was maintained at ~9 with aqueous ammonia. After 12 h, the precipitates were filtered, and washed with distilled water for three times, which was then cold-dried at ~50°C for 16 h to obtain MP powder. The surface morphology of MP powder was observed using scanning electron microscopy (SEM, S-4800N, Hitachi, Japan).

PS and WP were purchased from Shanghai Showa Highpolymer Co. Ltd (Shanghai, China) and Tokyo Chemical Industry Co. Ltd (Tokyo, Japan), respectively. PMWC scaffold with MP content of 50 wt%, PS content of 40 wt% and WP content of 10 wt% were fabricated by a method of solution casting-particle leaching. Briefly, PS particles and WP powders were dissolved into dimethyl formamide (DMF). MP was added into the solution, followed by continuous stirring to disperse uniformly. Sodium chloride (NaCl) particulates with diameters ranging from 300–500 μm were used as the porogens, which were added into the slurry and stirred for 30 min. The mixture was then cast into the molds (Φ12×2 mm and Φ
6×6 mm). The obtained samples were dried at 37°C for 48 h to remove the solvent. Afterwards, the samples were immersed into water for 48 h at 37°C to leach out the NaCl particulates (deionized water refreshed every 12 h), and then the samples were dried at 37°C for 12 h to obtain the PMWC scaffold. The PMC containing 50 wt% MP and 50 wt% PS, and PS scaffolds were prepared by the same method.

The scaffolds were characterized by X-ray diffraction (Geigerflex, Rigaku Co. Ltd, Japan) and Fourier transform infrared spectrometry (Magna-IR 550, Nicolet, East Rutherford, NJ, USA). The surface morphology and microstructure of the scaffolds were observed by scanning electron microscope (GeminiSEM 500, Gemini, Germany). The compressive strength of scaffolds was performed using a mechanical testing machine (HY-0230, Shanghai, China). Cylindrical scaffolds (φ 6×6 mm) were loaded with a 1 kN load at the speed of 1 mm/min until crushed. The compressive strength was determined from the peak of the stress-strain curve. Five samples of each group were measured and the average value of compressive strength was obtained. The porosity of the scaffolds was determined by using Archimedes method, which was calculated as following:

Porosity (%) = \( \frac{(m_2 - m_1)}{(m_2 - m_3)} \times 100\%

Where m1, m2 and m3 were the dry weight, wet weight and weight of the scaffolds suspended in water (subtracting buoyancy from m1), respectively.

**In vitro Bioactivity of Scaffolds**

In vitro bioactivity (apatite mineralization) of the scaffolds (φ12×2 mm) was evaluated by soaking the samples in SBF (pH=7.4) for different time periods. At each time point, samples were removed and the concentrations of Mg, Ca and P ions in SBF solution were determined by inductively coupled plasma atomic emission spectrometer (IRIS 1000, Thermo Elemental, Waltham, MA, USA). The samples were removed from the solution and rinsed with deionized water, and then dried in oven at 37°C for 24 h. The surface of the samples after soaking for 5 and 10 days was characterized by energy dispersive spectrometry (Falcon, Franklin Lake, NJ, USA) and SEM, respectively.

**In vitro Degradability of the Scaffolds**

The in vitro degradability of the scaffolds (φ12×2 mm) was assessed by measuring the weight loss of the samples after soaked into PBS for different time. In brief, the samples were weighed (W1) and immersed into PBS (pH=7.4) in sealed polyethylene bottles with a solid/liquid mass ratio of 0.1 g/20 mL. Each group was tested and incubated at 37°C in triplicate with constant shaking at 100 rpm for 12 weeks. The PBS solution was refreshed once a week. At different time points, the samples were removed from the solution and rinsed gently with deionized water. The samples were dried at 37°C until the water volatilized completely, and then weighed (W2). The weight loss was calculated according to the following equation:

\[
\text{Weight loss} = \left( \frac{W_1 - W_2}{W_1} \right) \times 100\%
\]

The pH value of the PBS solution containing the scaffolds was measured using a pH meter (FE20K, Mettler Toledo, Switzerland) at different time points.

**Cell Culture on the Scaffolds**

The MC3T3-E1 cells were purchased commercially from the Cell Bank of Type Culture Collection of Chinese Academy of Sciences, Shanghai, China. The cells were cultured in DMEM (Hyclone, Thermo Fisher Scientific, Waltham, MA, USA) supplemented with 10% FBS (Thermo Fisher Scientific, Waltham, MA, USA), 1% penicillin (100 U/mL) and streptomycin sulphate (100 μg/mL) (Thermo Fisher Scientific). The cells were cultured at 37°C in a humidified atmosphere of 5% CO2 in air, and the culture medium was replaced every other day.

**Cell Morphology**

After incubation for 24 h, the specimens were washed gently with PBS to remove the unattached cells. Then the cells on specimens were fixed with 4% paraformaldehyde for 15 min at room temperature and were permeabilized with 0.1% Triton X-100 in PBS for 10 min. After washed with PBS (×3), the cells were stained with DAPI (Sigma) and FITC (Sigma). The cell morphology was visualized using confocal laser scanning microscope (CLSM, Nikon A1R, Nikon, Japan).

**Cell Proliferation and Alkaline Phosphatase (ALP) Activity**

The cell proliferation was determined using the CCK-8 assay at one, three, and seven days postseeding according to a standard procedure. In brief, prior to culture with cells, the scaffold specimens (φ12×2 mm) were sonicated in ethanol for 24 h and sterilized under ultraviolet light for
24 h. MC3T3-E1 cells with a density of $1 \times 10^5$ cells/well were seeded on the samples, in 24-well tissue culture plates. At each time point, the samples were taken out, gently rinsed with sterilized PBS three times, and then transferred to a new 24-well plate. A total 500 μL of cell medium containing 50 μL of CCK-8 solution (Dojindo Molecular Technologies Inc., Kumamoto, Japan) was added to each well and empty wells containing DMEM were set up as a negative control. After three hours, the culture medium was collected and 100 μL of the supernatant was transferred into a 96-well plate and read at 450 nm using a microplate reader (Synergy HT, Bio-tek, Winooski, VT, USA) with 620 nm as a reference wavelength.

To evaluate the ALP activity, the samples were cultured with MC3T3-E1 cells with density of $2 \times 10^5$ cells/well in 24-well culture plates at 37°C with 5% CO$_2$ in an incubator for 7, 10 and 14 days. After incubation for 24 h, the culture medium was changed to the osteogenic induction medium, the minimum essential medium alpha modified (α-MEM) culture medium, which was supplemented with 10% FBS, 0.1 μM dexamethasone (Sigma), 50 mg/mL L-ascorbic acid (Sigma), and 10 μM β-glycerophosphate sodium (Sigma). These media were renewed every other day during two weeks. After 7, 10, and 14 days, the scaffolds were washed with PBS ($\times3$), and then lysed in a 0.2% Triton X-100 solution through four standard freeze-thaw cycles. The cell lysates were then mixed with 50 μL of p-nitrophenylphosphate (1 mM, Sigma, USA) substrate solution (pH=9) composed of 0.1 mol/L glycine and 0.5 mmol/L MgCl$_2$ in 1 M diethanolamine buffer and incubated at 37°C for another 15 min. Then the reaction was terminated by adding 50 μL of NaOH (1 M) solution. Then a spectrophotometer was used to determine the p-nitrophenol production in the reaction solution at wavelength of 405 nm. While the bicinchoninic acid (BCA) protein assay kit measured the total protein quantity for normalization.

**Animal Experiments**

This study was performed according to a protocol approved by the Research Center for Laboratory Animals of Shanghai Jiao Tong University Affiliated Sixth People’s Hospital, and followed with nation standard GB/T 35,892–2018 “Laboratory animal—Guideline for ethical review of animal welfare”. The 24 New Zealand white rabbits (male, ~2.0 kg) were chosen for surgery. Systemic anesthesia was achieved with 30 mg kg$^{-1}$ pentobarbital sodium and local infiltration with articaine with 1:100,000 noradrenaline. A median incision was made on the left leg of the rabbit, and then a 6 mm radial defect was introduced in the thighbone of the rabbits. The bone defects were filled with the scaffolds with the size of $\Phi 6 \times 6$ mm. Penicillin (4000 units per body) and streptomycin (200 mg per body) were administrated subcutaneously for prevention of infection. After scaffolds were implanted for 4, 8, and 12 weeks, rabbits were sacrificed with an overdose of intravenous pentobarbital solution and bone samples were collected and placed into 4% neutral buffered formalin.

**3-D SRmCT Imaging**

In order to evaluate the process of bone defects repair after the implantation, synchrotron radiation micro computed tomography (SRmCT) measurements were performed at beam line BL13W of Shanghai Synchrotron Radiation Facility (SSRF, Shanghai, China) using a monochromatic beam with an energy of 30 keV and a sample-to-detector distance of 1.5 m. In the current study, a 4000×2500 CCD detector with the pixel size set to 6 μm was used to record images. 1500 projections within an angular range of 180° were taken and the exposure time amounted to four seconds per projection. 2-D images were taken within a view of 6 mm in height and the samples were scanned from the top to the bottom successively. Finally, the 2-D images of the whole sample were integrated by Adobe Photoshop CS5 Software (Adobe Systems Incorporated, San Jose, CA, USA). 3-D images of the samples were reconstructed using a filtered backprojection algorithm. The images were finally converted to an eight-bit data format, proportional to the measured attenuation coefficients of the voxels. The VG Studio MAX 2.0 software (Volume Graphics, Heidelberg, Germany) served for the visualization of the tomographic data.

**Histological Evaluation**

After SRmCT scanning, the bone samples were decalcified in 10% EDTA (pH=7.0) for around 30 days at room temperature, and then washed with double-distilled water gently, dehydrated in ascending grades of alcohol, and embedded in paraffin. Serial section of the samples was made into 20 μm (thickness) by a microtome and then frozen at $-20°C$. One section was selected out of 10 sections of each sample were performed with H&E staining. The new bone formation within scaffolds was observed under a light microscope (TE2000U, Nikon, Japan).
The histological sections were used to quantitatively determine the amount of newly formed bone after the samples implanted in vivo for different times. Three pieces of histological sections were randomly selected from the PS, PMC, and PMWC. After H&E staining, each section was observed under light microscope, and at least six images were randomly obtained from the same section. Using image analytical software Image-ProPlus (Media Cybernetics, Rockville, MD, USA), new bone area was expressed as percentage of newly formed bone area within the original bone defect area, after the scaffolds implanted for 4, 8, and 12 weeks.

**Immunohistochemical Analysis**

Decalciﬁed histological sections were mounted on amino-propyl-triethoxy-silane coated slides. These sections were then dewaxed in descending concentrations of alcohol and rehydrated. The slides were immersed into 3% hydrogen peroxide to block endogenous peroxi-dases and rinsed with PBS. Then the slides were immersed into 0.1% Triton X100 in PBS for 20 min to allow penetration of the membrane. Antigen retrieval was carried out in a 10 mM warm citrate buffer at 60°C for 15 min at pH=6.0. Afterwards, the slides were rinsed gently with PBS. Speciﬁc sites were saturated with normal goat serum for 40 min at 37°C. The sections were incubated with the speciﬁc antisera (all diluted 1:100) overnight at 4°C.

The histological sections were deparafﬁnized and incubated with primary antibodies against BMP-2 (1:25 dilution; Abcam, Inc., Cambridge, UK) to identify osteogenesis before applying ﬂuorescent donkey polyclonal secondary antibody IgG (diluted 1:200, Invitrogen) for 30 min at room temperature. Thereafter, the sections were incubated for 30 min with DAPI for counterstain. The slides were then examined microscopically (TE2000U, Nikon, Japan).

Alternatively, deparafﬁnized sections were incubated with primary antibodies against VEGF (1:100 dilution, Abcam) to identify blood vessels, following incubation with a secondary antibody. Then DAB substrate (DAKO, Cambridge, UK) was used to stain the slides, and the slides were counterstained with hematoxylin and mounted. At last, slides were assessed by using a light microscope. Quantitatively analysis of the positive expressions of BMP-2 and VEGF was performed by using Image-ProPlus after the implantation of scaffolds at 4, 8, and 12 weeks.

**Statistical Analysis**

All quantitative data expressed as the mean ±SD were analyzed with Origin 8.0 (Origin Lab Corporation, USA). Statistical comparisons were carried out using analyses of one-way ANOVA followed by Tukey’s post hoc test. Statistical significance was attained with greater than 95%CI (p<0.05).

**Results**

**Characterizations of the Composite Scaffolds**

Figure 1A and B shows the scanning electron microscopy (SEM) images of MP powders at different magnifications. Clearly, the MP particles displayed flake-like shapes with an irregular size of ~500 nm Figure 1C–H shows the SEM images of the PS, PMC and PMWC scaffolds. Macroscopically, all scaffolds showed well interconnected macropores with a pore size ranging from 400 μm to 600 μm. The PS images at higher magnification (Figure 1D) showed a smooth surface. Conversely, PMC exhibited a slightly coarse surface because of the introduction of MP particles into PS. More interestingly, the PMWC not only possessed macropores, but also had micropores ranging from 10 μm to 20 μm on the walls of the macropores (Figure 1H).

The compositions of PS, PMC, PMWC, MP, and WP were conﬁrmed by X-ray diffraction (XRD) as shown in Figure 2A. The peaks at 2θ=19.3°, 22.2°, attributed to PS, suggested that PS had good crystallization, whereas the peaks at 2θ=12.2°, 13.5°, 21.1°, 23.2°, 29.6°, 31.4°, 34.7° and 36.4°, assigned to MP, demonstrated the weak crystallization of MP. The broad, instead of sharp, peak at around 2θ=20° in the XRD of WP indicated amorphous phase.

Figure 2B illustrates the Fourier transform infrared (FTIR) spectra of PS, PMC and PMWC, MP and WP. The characteristic band of PS at 1250–1000 cm\(^{-1}\) corresponded to the C=O stretching of the ester carbonyl group. The characteristic bands at 1746, 1308, 1197 and 1042 cm\(^{-1}\) of MP were assigned to the P-O stretching and bending. The characteristic band of WP at 1700–1600 cm\(^{-1}\) corresponded to the C=O–NH–stretching and bending. Evidently, the spectra of the PMC and PMWC exhibited the characteristic bands from PS and MP, and from PS, MP, and WP, respectively.

The results revealed that the compressive strengths of PS, PMC and PMWC scaffolds were 1.9 MPa, 2.8 MPa and 3.0 MPa, respectively. Moreover, the porosity of PS, PMC and PMWC scaffolds were 71.3%, 73.2% and 72.1%, respectively.
In vitro Bioactivity

The in vitro bioactivity (apatite mineralization) of scaffolds was investigated by soaking the samples in simulated body fluid (SBF) for different time as shown in Figure 3. SEM images of the scaffolds after immersed into SBF for 5 (Figure 3A, C, E) and 10 (Figure 3B, D, F) days were presented. PS, PMC and PMWC scaffolds showed different morphologies. Evidently, PS surface did not show
change (no apatite formation on scaffold surface) before and after immersion in SBF, indicating poor bioactivity of PS. In contrast, a layer of ball-like particles, the characteristic of apatite, formed in all composite scaffolds with the incorporation of MP after immersion in SBF. Clearly, the PMWC were fully covered with a new layer of apatite after 10 days, with bigger particle size and more uniform coverage than PMC, an evidence of faster apatite growth, meaning better bioactivity, on PMWC.

Figure 3G and H presented the energy-dispersive spectroscopy (EDS) of PMC and PMWC scaffolds after soaking into SBF for different times, respectively. At 5 or 10 days, P and Ca peaks appeared and increased visibly, indicating Ca-P-containing particles deposited on the scaffolds. This should be attributed to the precipitation of apatite layer on scaffolds. Compared with PMC, PMWC showed higher peak intensity, consistently, evidencing the better bioactivity of PMWC than PMC. These results demonstrated that apatite formation was induced on PMC, due to the addition of MP, and the WP in PMWC further promoted the apatite mineralization compared with PMC.

Ionic Concentration Changes of SBF Solution
Changes in the ion concentrations of SBF solution were investigated using inductively coupled plasma-atomic emission spectroscopy (ICP-AES) after the immersion of PMC and PMWC scaffolds for different time periods as shown in Figure 4. Both Mg and P ion concentrations in SBF rose with time, but at different rates, while Ca ion concentration in SBF declined with time. For PMC, at 10 days, the Mg, P, and Ca ions concentrations reached 75.3 mg L$^{-1}$, 52.6 mg L$^{-1}$, and 53.6 mg L$^{-1}$, respectively. Differently, the concentration of Mg, P and Ca ions in SBF solution soaking PMWC reached 92.0 mg L$^{-1}$, 65.0 mg L$^{-1}$, and 49.6 mg L$^{-1}$ at 10 days,
When the samples were immersed in SBF, Mg and P ions released from scaffolds into SBF solution, while Ca and P ions in the solution deposited to form apatite on the material surfaces, and the release of P was the leading process, in agreement with the EDS analysis in Figure 3G and H.

**Figure 3** SEM images of surface morphology of PS (A, B), PMC (C, D), and PMWC (E, F) immersed into SBF for 5 (A, C, E) and 10 (B, D, F) days. EDS of PMC (G) and PMWC (H) immersed in SBF for 5 and 10 days, respectively.

**Abbreviations:** SEM, scanning electron microscope; PS, polybutylene succinate; PMC, polybutylene succinate-magnesium phosphate composite scaffolds; PMW, polybutylene succinate-magnesium phosphate-wheat protein composite scaffolds; SBF, simulated body fluid.
In vitro Degradability and pH Change in PBS

As a great candidate scaffold for bone tissue repair, the degradation kinetic was expected to synchronize the newly formed bone tissue. Hence, the evaluation of degradation of biomaterials was of importance prior to further clinical application. Figure 2C shows the weight loss of the scaffolds after immersion into PBS for different time intervals. Encouragingly, the weight loss of PMWC scaffolds increased with the time and reached 58.43 w% at 12 weeks, while PMC and PS were 46.95 w% and 9.98 wt %, respectively. The results indicated that the incorporation of WP into PMWC improved the degradation kinetics, inducing a faster degradation rate than PMC.

The pH change in solution with time after the samples soaked into PBS for different time as shown in Figure 2D. It was found that the pH value corresponding to PMWC increased from initial 7.40 to 7.51 at 10 days and corresponding to PMC increased from initial 7.40 to 7.59, whereas the pH for PS reduced from initial 7.40 to 7.32. Significantly, the addition of MP into PS efficiently increased the alkaline condition, which deviated from the normal physical pH of 7.4 too much. Interestingly, the addition of WP into the PMC not only buffered the increase of pH value during degradation, slightly higher than physical pH, but also inhibited the occurrence of acidity, as observed in PS degradation.

Cell Morphology, Proliferation and Differentiation

Figure 5 illustrates the confocal laser scanning microscopy (CLSM) images of the MC3T3-E1 cell growth on the scaffolds stained by fluorescein isothiocyanate (FITC) and DAPI, at 1, 3 and 7 days. The cells exhibited a typical fibroblastic morphology with more cytoplasmic extensions and better filopodial attachments on PMC than PS, and PMWC scaffolds showed even better cell spreading than PMC. Alternatively, the cells showed denser growth on composite scaffolds, especially, PMWC, as compared with PS.

Figure 6A showed the proliferation of MC3T3-E1 cells on scaffolds measured by the cell counting kit-8 (CCK-8) assay. The relative proliferation rates for these scaffolds in tissue culture plate (TCP) increased with culture time, meaning that the cell numbers on scaffolds were sustainably increased. Notably, the cells on PMWC exhibited a prominently higher proliferation rate than both PMC and PS at seven days. However, at three and seven days, no statistical difference was found among PS, PMC and PMWC scaffolds at one and three days. Figure 6B shows the alkaline phosphatase (ALP) activity of MC3T3-E1 cells on scaffolds at 7, 10, and 14 days. The MC3T3-E1 cells on PS showed lower ALP activity than those cultured on PMWC and PMC at all the time points.

Macro Observation and SRmCT Analysis

The photos of the bone defects implanted with PS, PMC, and PMWC scaffolds for 4, 8, and 12 weeks were shown in Figure 7. Visibly, the bone defects were gradually healed with time. After 12 weeks, more significantly, the bone defects implanted with PMWC were replaced with newly formed bone tissue, whereas they were not completely renovated for PS.

The bone samples after implantation in vivo for 4, 8, and 12 weeks were scanned in SRmCT system to evaluate the in vivo osteogenesis of the scaffold (Figure 8). At four weeks, the 3-D images from SRmCT revealed that obvious new bone formation in PMWC scaffolds.
occurred throughout the cross section of the defects. Conversely, smaller amount of new bone in PMC was recognized at the native bone margins and the defect periphery. In addition, the PS exhibited bone formation at the outer surface of the cylinder, however, inferior compared with PMC, especially PMWC. This difference among PS, PMC, and PMWC was also found at eight weeks, and the bone repair was slightly improved compared with four weeks. At 12 weeks, the most extensive bone ingrowth, along with the formation of new bone cortex, was observed throughout PMWC. The new bone area for PMC increased with time and was lower than PMWC. This situation was even worse for PS, which showed significantly less bone accumulation with a large area of hole visible in the defect sites, and the bone defects, therefore, remained unrepaired.

**Histological Evaluation**

Figure 9A shows the images of histological sections, visualized by H&E staining, after PS, PMC, and PMWC scaffolds implanted into the femoral defects of rabbits for different time. After four weeks postimplantation, a few new bone tissues were found in PMC and PMWC and increased gradually with implantation time. At 12 weeks, a large amount of new bone tissues were observed in the PMWC. New bone tissues were also found in PMC, but the area was smaller than that in PMWC. On the contrary, the new bone tissues in PS were less than PMC and PMWC from 4 to 12 weeks postimplantation.

Figure 9B shows the quantitative analysis of the new bone area in PS, PMC, and PMWC scaffolds at different time points. Consistently, the new bone area increased with the implantation time for all scaffolds, but the bone

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**Figure 5** CLSM images of MC3T3-E1 cells stained by FITC (green) and DAPI (blue) after cultured on PS, PMC, and PMWC for 1, 3, and 7 days. Scale bar of 50 μm is applicable to all.

**Abbreviations:** CLSM, confocal laser scanning microscopy; MC3T3-E1, mouse pre-osteoblastic cell line; FITC, fluorescein isothiocyanate; PS, polybutylene succinate; PMC, polybutylene succinate magnesium phosphate composite scaffolds; PMWC, polybutylene succinate magnesium phosphate-wheat protein composite scaffolds.
regeneration in PS was extremely sluggish from 4 to 12 weeks, relative to other groups. At four weeks, there was no significant difference between the PMC and PMWC. However, at 8 and 12 weeks, the new bone areas for PMWC (8 w:55.46%, 12 w:90.87%) were prominently higher than PMC (8 w:43.77%, 12 w:75.35%) and PS (8 w:8.12% and 12 w:10.27%).

Immunohistochemical Analysis
Figure 10 shows the immunohistochemical staining of bone morphogenetic protein-2 (BMP-2, Figure 10A) and vascular endothelial growth factor (VEGF, Figure 10C) for PS, PMC, and PMWC scaffolds after implanted in vivo for 4, 8, and 12 weeks. Overall, the BMP-2 staining area (brown) increased with implantation time, indicating that...
Figure 8 3-D reconstruction images of bone samples from SRmCT after PS, PMC, and PMWC implanted into bone defects of rabbit femur for 4, 8, and 12 weeks. 
Abbreviations: SRmCT, synchrotron radiation micro-computed tomography; PS, polybutylene succinate; PMC, polybutylene succinate-magnesium phosphate composite scaffolds; PMWC, polybutylene succinate-magnesium phosphate-wheat protein composite scaffolds.

Figure 9 (A) Histological evaluation (H&E staining) of new bone formation and materials degradation after PS, PMC, and PMWC implanted in vivo for 4, 8, and 12 weeks. (B) Quantitative analysis of the new bone area by histological sections after PS, PMC, and PMWC implanted in vivo for 4, 8 and 12 weeks. *p<0.05 and **p<0.01 relative to Ps. 
Abbreviations: PS, polybutylene succinate; PMC, polybutylene succinate-magnesium phosphate composite scaffolds; PMWC, polybutylene succinate-magnesium phosphate-wheat protein composite scaffolds.
the positive expressions of BMP-2 gradually increased with time. At 12 weeks, the positive expression of BMP-2 on PMWC was much higher than on PMC, indicating that the osteogenic properties of PMWC were better than PMC. Nevertheless, the PS did not show good osteogenic properties from 4 to 12 weeks. In addition, the VEGF staining area (brown) increased with implantation time, indicating that the positive expressions of VEGF increased with time. At 12 weeks, the positive expression of VEGF for PMWC was higher than PMC, indicating that the vascularization of PMWC was better than PMC, while the PS showed the poorest vascularization from 4 to 12 weeks.

**Figure 10B** shows the quantitative analysis of positive expression of BMP-2 for PS, PMC, and PMWC scaffolds, which increased with implantation time. In a good agreement with **Figure 10A**, the positive expressions for PMWC were 20.66%, 41.12%, and 73.28%, at 4, 8, and 12 weeks, respectively. These values decreased to 17.24%, 34.06%, and 58.67% for PMC and 6.23%, 12.29%, and 21.77% for PS, respectively. **Figure 10D** shows the quantitative analysis of positive expression of VEGF for PS, PMC, and PMWC scaffolds, which increased with implantation time. The positive expressions of VEGF for PMWC were 16.62%, 32.15%, and 59.28%, and slightly decreased to 14.27%, 24.03%, and 42.69% for PMC, whereas they
dramatically dropped to 3.27%, 8.19%, and 15.71% for PS at 4, 8, and 12 weeks, respectively.

Discussion
PS possesses good biocompatibility, degradability, flexibility and processability. However, PS does not meet the requirement of bone regeneration due to its disadvantages, such as biological inactivity and slow degradation rate, etc. Natural polymers are the most important raw materials for the development of biomedical materials. As one of the natural polymers, WP with excellent properties, such as good biocompatibility and biodegradability, has been applied in many fields including coating, encapsulation, drug release, and other biomedical fields. In this study, PMWC (containing 50 wt% PS, 40 wt% MP and 10 wt% WP) scaffold was fabricated using a method of solution casting and particulate leaching. The as-fabricated PMWC scaffold exhibited the well interconnected macroporous structure with the pore size ranging from 400 μm to 600 μm, which were consistent with the size distribution of NaCl particles used as porogens. More interestingly, the PMWC not only possessed macropores, but also had micropores ranging from 10 μm to 20 μm on the walls of the macropores. The PMC and PMWC scaffolds showed slightly coarse surface because of the introduction of MP or MP and WP into PS while the PS exhibited smooth surface.

The compressive strengths of PS, PMC and PMWC scaffolds were 1.9 MPa, 2.8 MPa and 3.0 MPa, respectively. The results demonstrated that the addition of MP into PS improved compressive strength of the composite scaffolds, and the addition of 10 wt% WP into PS had no obvious effects on the compressive strength of PMWC. The porosity of PS, PMC and PMWC scaffolds were 71.3%, 73.2% and 72.1%, respectively. The results suggested that the incorporation of MP into PS did not have obvious effects on the porosity of the scaffolds as compared with PS. Moreover, PS had no obvious effects on the porosity of PMWC compared with PMC.

The ability to induce the apatite formation is an indicator of generating interfacial bonding with bone tissue when implanted in vivo. Apatite formation on the biomaterial surface when interacting with relevant biological fluids, such as SBF here, can verify in vitro bioactivity and predict the in vivo bone regeneration ability. The apatite formed efficiently on PMWC and PMC surfaces after immersion in SBF, whereas no apatite formation on PS, indicating that the bio-inert PS had been converted into bioactive scaffolds by incorporating MP and WP into PS. In addition, much more apatite formed on PMWC than PMC, suggesting that the WP exerted a positive influence on promoting apatite formation on PMWC. The presence of WP might have some effect on the microstructure, wettability or degradability of PMWC scaffolds, thus promoting the apatite mineralization. The apatite mineralization on PMWC and PMC surfaces could be interpreted as: once the degradation of MP was triggered in SBF, Mg and P ions released into the solution, meanwhile, the Ca and P ions in the solution precipitated on the scaffolds to form apatite, in which P ion release was considered as the leading process, which was consistent with the increase of Mg and P ion concentration, and the decrease of Ca ion concentration. The change of ion concentration in solution for PMWC was more significant than that of PMC because of the fast degradation of WP in the composites.

In the design and manufacture of scaffolds for bone tissue regeneration, appropriate degradability is desired. In this study, only a slight weight loss was observed for PS during the entire in vitro degradation period due to a random and bulk hydrolysis of the ester bonds in the PS chain. However, the weight loss of PMC and PMWC were more significant than PS, and gradually increased with time due to the degradation of MP in the composites. Furthermore, the weight loss of PMWC was higher than PMC because PMWC contained WP with a fast degradation rate. Meanwhile, the micropores in PMC and especially PMWC instead of smooth surface in PS, which might enlarge the contact area of scaffolds with PBS, improving degradation rate of PMWC in particular, which was in good agreement with previous studies.

The release of acidic degradation by-products from some biomaterials would lead to inflammatory responses in vivo and many approaches have been tried to control the pH decrease during the degradation of biomaterials. In this study, the pH value for PS decreased from 7.40 to 7.32, indicating the presence of an acidic product during degradation. However, the pH for PMC showed slight increase from the initial 7.40 to 7.59 during the degradation, this might be attributed to the dissolution of the MP, which released alkaline ions (Mg2+). Furthermore, the pH for PMWC showed more moderate increase from the initial 7.40 to 7.51 during the degradation, indicating that some acidic products, such as amino acids, might be produced from WP degradation. Previous study has shown that weak alkaline condition provided a better microenvironment for
cell proliferation and subsequently differentiation. Therefore, PMWC might supply a good microenvironment for cell proliferation and cell ingrowth both in vitro and in vivo.

The in vitro cell culture demonstrated that the cells on both PMWC and PMC presented better spreading with more pseudopods anchoring on the scaffold surface than on PS. This could be understood in two aspects: firstly, the presence of micropores on the walls of macropores of PMWC and PMC scaffolds, rather than PS, might be conducive to pseudopod anchorage, as suggested in earlier reports. Secondly, MP and WP in the composites might have obvious effects on the cell growth, evidenced by the fact that the cell morphology on PMWC were better than PMC because of the incorporation of WP, and much better than PS due to the incorporation of both MP and WP. The subsequent cell proliferation and ALP results showed a similar tendency. Previous studies have shown that the Mg dissolution products from bioactive glasses/ceramics stimulated proliferation and differentiation of osteoblasts. Apparently, herein, the addition of WP improved the degradation rate of PMWC compared with PMC because of the fast degradation of WP. Furthermore, the fast degradation of PMWC promoted the degradation of MP in the composites, which further accelerated the Mg ion release from PMWC, and thus promoted cell proliferation and differentiation.

Implantation of PMWC scaffolds into femur defects of rabbits for different time periods was performed for a further investigation on in vivo osteogenesis and biocompatibility. The SRmCT results demonstrated that the bone defects were gradually repaired by PMWC, significantly faster than PMC and PS, indicating that the PMWC promoted the restoration of bone defects. The histomorphometry analysis showed that the new bone tissues in both PMWC and PMC increased with time, indicating good osteogenesis. In addition, at both 4 and 12 weeks, more new bone tissues were found in PMC than in PS, indicating that the addition of MP into PS promoted the new bone formation in PMC. Furthermore, more new bone tissues were found in PMWC than in PMC, indicating that the addition of WP into PMC further improved the new bone regeneration in PMWC. Our results demonstrated that PMWC containing MP and WP stimulated bone tissue growth more efficiently in terms of the healing rate and quality of bone regeneration.

BMP-2 has been demonstrated to potently induce osteoblast differentiation in a variety of cell types. In addition, the BMP-2, like other bone morphogenetic proteins, plays an important role in the development of new bone, and the positive expression of BMP-2 is known to be of major importance in bone generation and reconstruction. Moreover, the vascularization of bone graft and bone substitute is the guarantee of successful transplantation. The procedure of vascularization involves the expression of VEGF, which has been identified as the key regulator factor for new bone regeneration. In this study, to evaluate the osteogenesis and vascularization of the scaffolds, the immunohistochemical staining and quantitative analysis of positive expression of BMP-2 and VEGF for PS, PMC, and PMWC scaffolds were carried out after being implanted in vivo for 4, 8, and 12 weeks.

The positive expression of BMP-2 for PMWC increased with implanted time, indicating that the osteogenic property of PMWC in vivo increased with time. In addition, the positive expression of BMP-2 for PMWC were higher than PMC, and PMC were higher than PS, indicating that the addition of MP into PS improved osteogenic property of PMC, and addition of WP into PMC further improved the osteogenic property of PMWC. Moreover, the positive expression of VEGF for PMWC increased with implanted time, indicating that the vascularization of PMWC increased with time. Furthermore, the positive expression of VEGF for PMWC were higher than PMC, and PMC were higher than PS, indicating that the addition of MP into PS promoted vascularization in PMC, and addition of WP further improved the vascularization of PMWC. The immunohistochemical analysis further confirmed that the PMWC containing MP and WP could promote osteogenesis and vascularization. Briefly speaking, this study demonstrated that the interconnected micro-macroporous scaffold of PS/MP/WP ternary composite with improved bioactivity and degradability significantly promoted the osteogenesis and vascularization in vivo, which might have great potential to be applied for bone tissue generation and to repair bone defects.

Conclusion
The interconnected micro-macroporous scaffold of PS/MP/WP ternary composite was fabricated in this study. The results demonstrated that incorporation of MP into PS obviously improved the degradability and bioactivity of PMC scaffold. Moreover, addition of WP into PMC further enhanced the degradability and bioactivity of PMWC scaffold. The in vitro cell experiments demonstrated that the PMWC scaffold containing MPC and WP
significantly promoted the cells proliferation and differentiation as compared with PMC and PS. The in vivo study showed that the PMWC containing MP and WP remarkably stimulated the new bone formation and ingrowth compared with PMC and PS. Furthermore, the results of immunohistochemistry further offered the evidence that PMWC promoted osteogenesis and vascularization. In short, the PMWC exhibited good biocompatibility, bioactivity, degradability, osteogenesis and vascularization, which would be an excellent ternary composite scaffold for bone tissue regeneration.

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Disclosure
The authors report no conflicts of interest in this work.

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