Enhancing cartilage repair with optimized supramolecular hydrogel-based scaffold and pulsed electromagnetic field

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** ABSTRACT **

Functional tissue engineering strategies provide innovative approach for the repair and regeneration of damaged cartilage. Hydrogel is widely used because it could provide rapid defect filling and proper structure support, and is biocompatible for cell aggregation and matrix deposition. Efforts have been made to seek suitable scaffolds for cartilage tissue engineering. Here Alg-DA/Ac-β-CD/gelatin hydrogel was designed with the features of physical and chemical multiple crosslinking and self-healing properties. Gelation time, swelling ratio, biodegradability and biocompatibility of the hydrogels were systematically characterized, and the injectable self-healing adhesive hydrogel were demonstrated to exhibit ideal properties for cartilage repair. Furthermore, the new hydrogel design introduces a pre-gel state before photo-crosslinking, where increased viscosity and decreased fluidity allow the gel to remain in a semi-solid condition. This granted multiple administration routes to the hydrogels, which brings hydrogels the ability to adapt to complex clinical situations. Pulsed electromagnetic fields (PEMF) have been recognized as a promising solution to various health problems owing to their noninvasive properties and therapeutic potentials. PEMF treatment offers a better clinical outcome with fewer, if any, side effects, and widely used in musculoskeletal tissue repair. Thereby we propose PEMF as an effective biophysical stimulation to be 4th key element in cartilage tissue engineering. In this study, the as-prepared Alg-DA/Ac-β-CD/gelatin hydrogels were utilized in the rat osteochondral defect model, and the potential application of PEMF in cartilage tissue engineering were investigated. PEMF treatment were proven to enhance the quality of engineered chondrogenic constructs in vitro, and facilitate chondrogenesis and cartilage repair in vivo. All of the results suggested that with the injectable self-healing adhesive hydrogel and PEMF treatment, this newly proposed tissue engineering strategy revealed superior clinical potential for cartilage defect treatment.

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1. Introduction

Due to the avascular nature, cartilage lacks innate self-repair ability to restore physiological structure and functions. Therefore, cartilage defect resulting from sudden injury or gradual wear and tear will eventually progress to osteoarthritis over time [1]. Current available treatments are mainly prescribed for symptoms relief only. Functional tissue engineering strategy using biomaterials has been proposed and actively investigated to provide innovative approach for the repair/regeneration of damaged cartilage.

Variety of natural and synthetic materials have been used as potential carrier biomaterials of cells or therapeutic agents for cartilage repair [2]. Hydrogels made of various polymers have been shown to promote the chondrogenesis and cartilage regeneration in the presence of inductive factors by providing the conducive 3D microenvironment [3, 4].

Specially, hydrogels fabricated by natural polymers, such as hyaluronic acid (HA) [5–7], collagen [8], chitosan [9,10], and alginate [11], have been widely used as defect-filling treatments because of their biocompatibility, low immunogenicity, biodegradability, and suitable physical properties. Li et al. [7] prepared an in situ forming injectable hydrogel system composed of hyperbranched poly (ethylene glycol) (HB-PEG) and thiol-functionalized hyaluronic acid (HA-SH) crosslinked via thiol-ene reaction. The as-prepared hydrogels exhibited tunable and stable mechanical properties and rapid in situ gelation rate under physiological conditions. Besides, based on the HB-PEG, thiolated-chondroitin sulfate (CS-SH) was selected to fabricate CS-SH/HB-PEG hydrogels [12]. The prepared hydrogel exhibited rapid gelation, excellent mechanical properties and prolonged degradation properties.

In our previous work [13], an effective “Host-Guest Macromer” (HGM) approach for the preparation of supramolecular gelatin hydrogels was studied. The HGM hydrogels are self-healable, injectable, re-moldable and shows ideal mechanical properties due to the reversible nature of the host-guest interactions and chemically crosslinked acrylated structure. Herein, based on the previous study, we would like to introduce a modified hydrogel design by supplementing dopamine functionalized alginate (Alg-DA) to the original formula (photo-crosslinkable acrylated β-cyclodextrins (Ac-β-CDs) and gelatin). In this design, chemical and physical multiple crosslinking mechanisms were simultaneously incorporated into the hydrogel, including UV-induced covalent bonds, host-guest interactions, hydrogen bonding, and π-π aromatic stacking interactions (Fig. 1 a). Therefore, better physical properties could be achieved to better meet the complicated requirements in cartilage regeneration. The multi-crosslinked structure improves their injection abilities, mechanical performance, and better self-healing abilities, and further endows the hydrogel with new physical properties like suitable swelling ratio, appropriate degradation rate and tissue-adhesive ability. The new design introduces a pre-gel state during hydrogel preparation, where mixed solution exhibit increased viscosity, allowing the pre-gel to remain in a semi-solid condition (Figure S1 a).

While suitable scaffolds can be designed, they cannot create high-quality cartilage tissue independently. The complicated requirements in cartilage regeneration often led to the failure of cartilage repair [18]. MSCs based stem cell therapy and tissue engineering strategy represent a promising solution. MSCs can differentiate into numerous cell types—including chondrocytes, fibrochondrocytes, and hypertrophic chondrocytes—resulting in a mixture of cartilaginous, fibrous, and hypertrophic tissues [19,20]. The success of MSC-based techniques may
remain limited if the presence of fibrous and hypertrophic tissue cannot be eliminated. Therefore, biological augmentative strategies are urgently needed to improve the therapeutic outcomes of tissue engineering in cartilage repair. In orthopaedics field, previous studies have proposed “the diamond concept” [21], which embodies a 4th element, environmental biophysical stimulation, to underpin tissue engineering. As a safe, noninvasive biophysical strategy with fewer, if any side effects for tissue repair, pulsed electromagnetic fields (PEMF) could not only benefit cartilage-related cells in vitro including promoting proliferation, increasing anabolic activities, and antagonizing the catabolic effects of inflammation [22–25], but also provide chondro-protective and chondro-regenerative effects on articular cartilage in vivo [26–29]. Therefore, we purpose that with the promising chondrogenic effects both in vitro and in vivo, PEMF could improve the quality of engineered biomaterial in vitro, and enhance cartilage repair in vivo, making it suitable as the 4th element in cartilage tissue engineering via augmenting the benefit of biomaterials consists of cells, scaffolds and bioactive factors.

In this study, we hypothesized that PEMF signal could benefit MSCs-based cartilage tissue engineering and promote cartilage regeneration, and with Alg-DA/Ac-β-CD/gelatin hydrogel, we investigated the effects of PEMF stimulation on MSCs chondrogenesis in vitro and in vivo, and its effect on enhancing cartilage repair in combination of MSCs-based tissue engineering using a rat osteochondral defect model.

2. Methods

All experiments were approved by the Animal Research Ethics Committee of the Chinese University of Hong Kong (AEEC number:20-225-HMF).

2.1. Isolation of rat BMSCs and human BMSCs

Green fluorescent protein (GFP) Sprague-Dawley rats (male, 8 weeks, SD-Tg (CAG-EGFP) Cz-004Osb) were used for rat BMSCs isolation. rBMSCs were obtained from the bone marrow of GFP-SD rats by density gradient centrifugation (850 ×g, 20 min). Under the approval of IRB of author’s institution (CREC Ref No. 2014.291-T), human BMSCs (hBMSCs) from three patients with osteoarthritis (OA) and undergoing total knee replacement surgery were obtained as pre-seeded cells. The obtained cells were cultured in αMEM, Gibco, ThermoFisher Scientific, USA) supplemented with 10% fetal bovine serum (FBS, Gibco, ThermoFisher Scientific, USA) and 1% penicillin/streptomycin (PS, Gibco, ThermoFisher Scientific, USA) (500 cells/cm², 37 °C, 5% CO₂) to isolate BMSCs and form colonies; medium was changed every 3 days. Cells at passage 3–5 were used for all following experiments.

2.2. Hydrogel preparation

2.2.1. Materials

All the reagents including sodium alginate (Alg), dopamine hydrochloride (DA), 1-(3-Dimethylaminopropyl)-3-ethylcarbodimide hydrochloride (EDC-HCl), N-hydroxysuccinimide (NHS) and 2-hydroxypropyl-4-(2-hydroxyethylxy)-2-methylpropophenone (Irgacure 2959) were purchased from Shanghai Aladdin Biochemical Technology Co., Ltd. The water used in the experiments was purified using Millipore Milli-Q Ultrapure water systems.

2.2.2. Synthesis and characterization of dopamine-modified alginate (Alg-DA)

2.0 g of Sodium alginate (10.1 mmol, according to the repeating unit of alginate molecule) was added into 200 mL of Milli-Q water and stirred until fully dissolved. Then EDC-HCl (1.936 g, 10.1 mmol) and NHS (1.162 g, 10.1 mmol) were added in the above alginate solution. The reaction solution was stirred for 60 min at room temperature. During this procedure, the carboxylic groups in the alginate molecules would be fully activated. After that, different amounts (0.639 g, 0.958 g, 1.915 g) of dopamine (3.37, 5.05, 10.1 mmol) were added into the above reaction solution and stirred for another 12 h at room temperature under N₂ atmosphere. Then, the reaction solution was dialyzed (MWCO 3500) against distilled water (pH 6) three days and the water was changed twice a day. Then the dialyzed materials solution was lyophilized. The corresponding product dopamine-modified alginate was denoted as Alg-DA-1, Alg-DA-2 and Alg-DA-3.

The successful preparation of synthesized Alg-DA was confirmed by nuclear magnetic resonance (¹H NMR) spectroscopy, which was performed through a Bruker 500 MHz NMR spectrometer (Bruker BioSpin GmbH). The samples were dissolved into D2O at a concentration of 10 mg/mL before the test.

The degree of dopamine substitution of different groups of Alg-DA was determined by measuring the absorbance at 280 nm using an ultraviolet-visible (UV-vis) spectrophotometer and calculating from a standard curve of dopamine hydrochloride solution. Different groups of Alg-DA solution were prepared with the concentration of 1 mg/mL and used as the tested samples. Dopamine hydrochloride solution was used to generate a standard curve with the concentrations ranging from 0.0156 to 1 mg/mL.

2.2.3. Synthesis of acrylated 1,4-cyclohexadien (Ac-β-CD)

The Ac-β-CD was synthesized and obtained according to our previous study [31].

2.2.4. Preparation of Alg/Ac-β-CD/gelatin and Alg-DA/Ac-β-CD/gelatin hydrogel

Firstly, gelatin was dissolved into dH₂O at 37 °C with fixed concentration of 5% (w/V). For the preparation of Alg/Ac-β-CD/gelatin hydrogel, 5 wt% of Ac-β-CD were added into gelatin solution under continuous stirring. After 20 min, 1% of Alg was added into the solution and thoroughly stirred. Then the initiator Irgacure 2959 was added at a concentration of 0.05% (w/V). The mixture was thoroughly stirred at 37 °C and a homogenous solution was obtained. The pH of the solution was kept at 7.2. Then the pre-gel solution was transferred into cell culture plate. Then 365 nm ultraviolet (UV) light (10 mW/cm²) was utilized to prepare the Alg/Ac-β-CD/gelatin hydrogels. For the preparation of Alg-DA/Ac-β-CD/gelatin hydrogel, the procedure was almost the same as above, except that the 1 wt% of Alg-DA was added instead of Alg.

2.3. Hydrogel swelling analysis

To evaluate the hydrogel swelling property, the prepared different groups of Alg/Ac-β-CD/gelatin hydrogels and Alg-DA/Ac-β-CD/gelatin hydrogels (500 µL of pre-gel solution for one sample) were lyophilized and then weighed to determine the dry weight of hydrogel material (W₀). Then the samples were incubated in PBS at 37 °C for 24 h to determine the water swelling degree. At different time points, the tests were taken out and the residual PBS on the sample surface was carefully removed, and the weight of the samples was recorded (Wₜ). After the test, the swelling ratio (SR) was calculated as the ratio of the weight of water in the swollen hydrogel at different time points to the weight of dry hydrogel: SR = (Wₜ - W₀)/W₀.

2.4. In vitro hydrogel degradation test

In vitro degradability of the Alg/Ac-β-CD/gelatin hydrogel and Alg-DA/Ac-β-CD/gelatin hydrogel was determined by weight changes of the hydrogels after incubated in PBS or Collagenase Type-I/PBS (0.1 mg/mL) at 37 °C for different time. The prepared different groups of Alg/Ac-β-CD/gelatin hydrogels and Alg-DA/Ac-β-CD/gelatin hydrogels (500 µL of pre-gel solution for one sample) were lyophilized and then weighed to determine the dry weight of hydrogel material (W₀). After that, the dried materials were incubated in PBS with 0.1 mg/mL of Collagenase Type-I for different time points. At the end of the test period, the weight of the hydrogel was recorded (Wₜ). The degradation rate (DR) was calculated as the ratio of weight loss to the initial dry weight: DR = (W₀ - Wₜ)/W₀.
hydrogel samples were immersed into PBS at 37 °C for 12 h for water equilibrium. Then the degradation test was conducted. The samples were removed from the incubation solution at different time of incubation. After carefully and thoroughly washing with distilled water, the samples were lyophilized and weighed (W1). Degradation rate (DR) of the hydrogel samples was calculated in terms of the following equation, DR = (W0 - Wt)/W0 x 100%.

2.5. Adhesion property

The adhesive properties of the as-prepared hydrogels were determined through the lap-shear adhesion measurements. By placing the hydrogel between two glass slides with the overlapped contact area dimensions of 25 mm x 20 mm x 0.5 mm, the samples were pressed slightly for 10 s to enhance the adhesion. Then the test samples were kept in 37 °C incubator for 12 h. Subsequently, the samples were pulled to failure and the max load was measured.

2.6. Induction of chondrogenic differentiation of MSCs-laden hydrogels and PEMF exposure in vitro

The therapeutic PEMF signal used in this study is designed by Orthofix Medical Inc. (Physio Signal, 101/s, 15 Hz, Orthofix Medical Inc, USA). As demonstrated in our previous study [32], our signal is safe and effective, and has been approved by FDA to manage long bone nonunions [33]. To investigate the effects of PEMF signal on chondrogenic differentiation of BMSCs in 3D culture environment, rBMSCs-laden hydrogels were divided into two groups. PEMF group received daily PEMF treatment 3 h/s/day for 21 days, and control group received no PEMF treatment. On day 0, rBMSCs-laden hydrogels were cultured in chondrogenic inductive medium (CIM, ThermoFisher Scientific, A1007101), and daily PEMF treatment (3 h/s/day) was given to the cells using an in vitro PEMF device (Orthofix Medical Inc.). Control group was kept under the same conditions without PEMF exposure. Samples were collected on day 7, 14, and 21 (n = 6 per group per time point), cut into two equal parts. One half was subjected to RNA extraction for quantitative real-time PCR analysis (n = 4/group/time point.), and the other half was fixed with 4% paraformaldehyde (PFA) following alcin blue, safranin O and immunohistochemistry staining (n = 4/group/time point.).

2.7. Nude mice transplantation

rBMSCs-laden hydrogels prepared as previous described were cultured in CIM for 14 days. Nude mice (male, 8 weeks old) were randomly divided into two groups: Hydrogel and hydrogel + PEMF. Hydrogels were transplanted subcutaneously into the back of nude mice according to the grouping, and daily PEMF treatment (3 h/s/day) were given to hydrogel + PEMF group via a specialized in vivo PEMF device (Orthofix Medical Inc.) [34]. The hydrogel group received no PEMF treatment. After 14 days of PEMF exposure, fresh samples were harvested for the mechanical test (n = 8). Other samples were cut into two halves. One half was subjected to RNA extraction for quantitative real-time PCR analysis (n = 8), and the other was fixed with 4% PFA following alcin blue, safranin O and immunohistochemistry staining (n = 8).

2.8. RNA-sequencing and bio-informatic analysis

hBMSCs were cultured in αMEM containing 10% fetal bovine serum and 100 U/mL penicillin with or without daily PEMF treatments (3 h/s/ day) for 7 days. Samples were then collected, and total RNA were extracted using RNeasy Mini Kit (Qiagen, Hilden, Germany). Total RNA was processed by mRNA enrichment method. The constructed DNA library was prepared as previously published [35], and RNA-sequencing was performed with BGISEQ-500 platform (Beijing Genomics Institute, Shenzhen, China) and hg38 genome database.

Differentially expressed genes (DEGs) were identified from RNA-seq data using an R package as previously described [36]. DEGs were defined as fold change ≥2 and Q-value ≤ 0.001. Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analysis was performed using DAVID (https://david.ncifcrf.gov). Candidate DEG functions and pathway interactions were analyzed using the ClueGO plug-in of Cytoscape.

2.9. Validation and pharmacological inhibition study of ERK and p38 pathways

After bioinformatics analysis and literature review, ERK and p38-MAPK pathways were located as possible pathways mediating the effects of PEMF on hBMSCs chondrogenesis. To further validate the involvement of ERK and p38-MAPK pathways on chondrogenic differentiation of hBMSCs during PEMF treatments, on day 0, cell pellets were cultured in CIM and received daily PEMF treatment (3 h/s/day) for 10 days. Cell pellets from control group were kept at same condition and received no PEMF treatment. Samples were collected on day 1, 3, 7 and 10 and subject to Western blot to analyze the expression and phosphorylation of ERK1/2 and p38 protein.

Pharmacological inhibition study was deployed to investigate the correlation between regulation of ERK and p-38 by PEMF treatment and chondrogenesis of hBMSCs. On day 0, Specific inhibitors of MEK / ERK (PD98059, 10 μmol/L, Invitrogen™, ThermoFisher Scientific, USA), p38 (SB20350, 3 μmol/L, Sigma-Aldrich, USA), or a combination of both inhibitors were added into CIM of pellet culture, and cell pellets received daily PEMF treatment (3 h/s/day) for 14 days. Samples were collected on day 1 to exam ERK1/2 and p38 activities after pharmacological inhibition through Western blot, and collected on day 7 and 14 for qPCR analysis of chondrogenic marker (Sox9) and hypertrophic marker (RUNX2).

2.10. Implantation of hydrogel into rat model of osteochondral defect

Three-month-old SD rats (male, weight 250–300 g, n = 40) were divided into 4 groups (n = 10), and the detailed experimental groups are shown in Table 1.

In brief, rat osteochondral defect models were established as previously described [37], and according to different groups, the osteochondral defects were transplanted with three types of hydrogels: blank hydrogels with no cells implanted, rBMSCs-laden hydrogels, and rBMSCS-laden hydrogels (Cells treated with daily 3 h PEMF in vitro exposure for 14 days), respectively. The osteochondral defects were created using a 1.5 mm diameter dental drill in the center of the femoral trochlear groove with 1.5 mm in depth. The defect was rinsed with 0.9% saline and the debris was removed using a curette. The defects were implanted with different biomaterials according to Table 1. Briefly, the pre-gel solution of Alg/Ac-β-CD/gelatin with initiator Irgacure 2959 was injected into the defects. Then 365 nm ultraviolet UV light (10 mW/cm²) was utilized to initiate the gelation. Daily PEMF treatment of 3 h/day was given to the rats 24 h after surgery until termination. Rats were terminated after 8 weeks with pentobarbital overdose. The right femur of the rats was collected and were fixed in 10% buffered formalin.

| Group | Material implanted | PEMF treatment after implantation |
|-------|--------------------|----------------------------------|
| Blank | Blank hydrogels (no cells) | No PEMF treatment |
| MSCs | rBMSCs-laden hydrogels | No PEMF treatment |
| P-MSCS | P-EMF enhanced rBMSCs-laden hydrogels | No PEMF treatment |
| P-MSCS + PEMF | P-EMF enhanced rBMSCs-laden hydrogels | Daily PEMF treatment, 3 h/ day |
Samples were decalcified in a 10% solution of buffered ethylenediaminetetraacetic acid (EDTA, Sigma-Aldrich, USA). Histological and immunohistochemical staining were performed to determine the healing outcome of osteochondral defect.

2.11. In vivo PEMF exposure

For in vivo PEMF treatment, nude mice or rats were placed in plastic cages for PEMF exposure to eliminate interference by metal components and placed into the in vivo PEMF system (Orthofix Medical Inc.). Same PEMF signal used in vitro was applied to the animals. Daily PEMF treatment of 3 h/day was given to the animals as described above. Nude mice or rats placed under the same condition but without PEMF exposure served as the control group.

2.12. Quantitative real-time PCR

Total cellular RNA was extracted from hydrogel samples with MiniBEST universal RNA extraction kit (Takara, Japan), and reversely transcribed into cDNA with M-MLV reverse transcriptase according to the manufacturer instructions (Takara, Japan). Real-time PCR was performed using the Step One Plus Real-Time PCR System (Applied Biosystems, USA). 10 μl reaction volume was consisted of 1 μl cDNA template diluted with Milli-Q water, 5 μl SYBR-Green Master Mix (2X, Applied Biosystems, USA), 3.4 μl PCR grade water, and 0.6 μl of primer (10 μM). The amplification procedure was carried out as follows: first at 95 °C for 5 min, and then 40 cycles of 95 °C for 15 s and 60 °C for 60 s. Primer sequences were shown in Table 2. The relative quantitative expression of gene was analyzed with the values of $2^{-\Delta\Delta C_T}$, normalized to the GAPDH expression.

2.13. Western blot

The cell pellets were collected and lysed using 50 μl radioimmunoprecipitation assay (RIPA) lysis buffer (Sigma-Aldrich, USA) with 1 mM phenylmethylsulfonyl fluoride (Roche, USA) at indicated time points. The lysate was centrifuged at 14 000 g for 15 min, supernatant collected and quantified using BCA assay kit (Thermo Fisher Scientific, USA). Normalized proteins were subject electrophoresis and electroblotted and probed with primary antibodies include ERK1/2 monoclonal antibody (1:1000, 13–6200, Sigma-Aldrich, USA), rabbit monoclonal anti-p-ERK1/2 (1:3000, ab184699, Abcam, UK), rabbit monoclonal anti-p38 (1:1000, ab70099, Abcam, UK), rabbit monoclonal anti-p–p38 (1:1000, ab4822, Abcam, UK) and mouse monoclonal anti-p–actin (1:1000, ab8226, Abcam, UK), overnight at 4 °C. Blots were then incubated with horseradish peroxidase (HRP)- conjugated secondary antibody (diluted 1:5000; Santa Cruz, CA, USA) at 37 °C for 1 h, and visualized using an ECL system (Millipore, Billerica, MA, USA). Optical intensity of each band were analyzed using ImageJ, and band intensity ratio ($A_{p\text{-ERK}}/A_{\text{ERK}}$ and $A_{p\text{-p38}}/A_{\text{p38}}$) were used for data analysis.

2.14. Mechanical test

Mechanical test was performed as previous described to obtain the compression Young’s modulus and ultimate strength [38]. Stress-relaxation analysis was conducted with MACH-1™ Mechanical Testing System (Biomomentum). The equilibrium Young’s modulus was calculated by the MACH-1 Analysis software (Biomomentum, Laval) with the obtained equilibrium load after 1000 s of relaxation at 50% strain under uniaxial and unconfined compression.

2.15. Safranin O & fast green staining

After deparaffinized and hydrated, sections of samples were stained with 0.05% fast green (Sigma-Aldrich) solution for 5 min. Sections were rinsed with 1% acetic acid solution for 15 s and stained with 0.1% safranin O (Sigma- Aldrich) solution for 5 min. Residual dye was removed using 95% ethanol and absolute ethanol for 2 min each.

2.16. Histology and immunohistochemistry staining

After deparaffinization, H&E staining was performed on rat cartilage sections, and immunohistochemistry staining was performed on hydrogel sections and rat cartilage sections. Immunohistochemistry staining was performed using a standard protocol as previously reported [39]. Sections were incubated with primary antibodies to mouse anti-rat Collagen type II (Col II; ab54712, 1:100, Abcam, UK), rabbit anti-rat Collagen type X (Col X; ab58632, 1:200, Abcam, UK) or rabbit anti-rat MMP13 (MMP13; ab39012, 1:200, Abcam) overnight at 4 °C; a horseradish peroxidase-streptavidin detection system (Dako, USA) was used, followed by counterstaining with hematoxylin. ImageJ was used to analyze percentage of positive area in the hydrogel samples or cartilage defect sites. For hydrogel samples, five randomly selected pictures from each section taken at 400 × magnification were used for semi-quantitative analysis.

2.17. Cartilage repair analysis by the Wakitani scoring system

Rat cartilage sections of each defect at week 8 after hydrogel implantation were scored using H&E staining together with IHC staining of Col II to define the region of cartilage in newly formed surface tissue (n = 10 per group). Histology sections were blindly scored by three independent researchers based on a previously established scoring system by Wakitani et al. [40].

### Table 2

| Gene Name   | Forward (5'–3') | Reverse (5'–3') |
|-------------|----------------|----------------|
| GAPDH       |                 | CACCACCCCTCTCTGATGCTATC |
| Sox-9       | AGCCGAGAATATCCAGTCTGCTGCTG | TCCCTGAGCAAACGTTGAAA |
| Col 2a1     | ACACCAAGGAGAAGGGGACCAGT | CCGGACTGAGGTTAGGAT |
| ACAN        | TTGTGGACTCTCAGGGGGTACACTC | GTCTGACGGAGGCCCACCAAG |
| MMP13       | AGGGCTCTACAGAAAGGCTTTC | GAGCTGCTGTCGCAAGGTTC |
| Col 10a1    | ATATCGCTGGGATGCTCAGGTTGC | TCCAGGTTTCACTTGTCAGC |
| Runx2       | GAACCCAGAAGGACGACGAGC | AATGGGCGCTTAAATCTATG |
| Human primers |                     |                 |
| Sox-9       | TGGGACGCTGCTGAAGAGACTTCTC | ATCCGGTGGTGGTCCTTTG |
| Runx2       | GACAAGACCAAAGTAATCATGGAACAGCAG | GTAAGCGCTTGTTGTTAAATCTCGT |

GAPDH: glyceraldehyde-3-phosphate dehydrogenase, Sox-9: SRY (Sex-Determining Region Y)-box transcription factor 9, Col 2a1: Collagen type II alpha 1 chain, ACAN: Aggrecan, MMP13: Matrix metalloproteinase 13, Col 10a1: Collagen type X alpha 1 chain, Runx2: Runx-related transcription factor 2.
2.18. Statistical analysis

Data was presented as mean ± standard deviation. All the quantitative and semi-quantitative data were analyzed using SPSS 18.0 software for windows (SPSS, Chicago, IL, USA). Parameters were analyzed by analysis of variance (ANOVA). One-way ANOVA and Tukey multiple comparison was used for comparison among groups and two-way ANOVA and Tukey multiple comparison was used for comparison among groups and time factor. p < 0.05 was considered as statistically significant.

3. Results

3.1. Characterizations of Alg-DA and gelation of Alg/Ac-β-CD/gelatin and Alg-DA/Ac-β-CD/gelatin hydrogels

To fabricate the supramolecular Alg-DA/Ac-β-CD/gelatin hydrogels, Alg-DA was firstly prepared. The 1H NMR spectra of the synthesized Alg-DA was presented in Supplementary Fig. S3. The results clearly showed the peaks at δ2.7–3.2, which were attributed to the –CH2–CH2– groups of dopamine, and peaks at δ6.6–6.9 ppm, which were attributed to the phenyl groups of dopamine. The results indicated that dopamine was successfully grafted to the alginate. To determine the conjugation efficiency of the dopamine substitution reaction, UV-vis spectra were utilized. First, a standard curve of absorbance at 280 nm of a series of preset dopamine concentrations was obtained. And the absorbance of 1 mg/mL dopamine modified alginate was also measured. Through the standard curve, the concentration of catechol in the Alg-DA sample solution could be calculated. And the amount of alginate in the sample solution is a fixed value. Thus, 6.7%, 9.4 and 15.3% of the alginate molecule units of Alg-DA-1, -2 and -3 were conjugated with catechol according to this calculation.

The gelation time of each group of hydrogels was monitored by the inverted vial method. Under UV irradiation, the gelation time of each group of hydrogels was recorded when the pre-gel solution transferred to solid-like state. The gelation time of all the groups was about 15–20 min, and the substitution degree of dopamine had no obvious influence on the gelation time.

As our hydrogel design was successfully carried out, the hydrogels demonstrated ideal physical properties (Fig. 1, Fig. S1.b, Video 1), owning to the chemical and physical multiple crosslinking and interactions during gelation. Along with the enhanced physical properties, the Alg-DA/Ac-β-CD/gelatin hydrogels were injectable through a 23G needle, and self-healable and re-moldable after injection (Video 1). It is also noteworthy that our design introduced a pre-gel state. By simply mixing the Alg-DA with Ac-β-CD/gelatin solution, the physical interactions are readily formed, resulting in increased viscosity and limited fluidity of the pre-gel (Fig. S1a). Subsequent UV initiated polymerization of Ac-β-CD, resulting in the gelation of Alg-DA/Ac-β-CD/ gelatin hydrogels that are more stable, physically strong, and mechanically robust.

Supplementary video related to this article can be found at https://doi.org/10.1016/j.bioactmat.2022.10.010

3.2. Hydrogel swelling test

The hydrogel swelling properties of the as-prepared hydrogels were determined through measuring the water uptake capacity. The swelling ratios of the Alg/Ac-β-CD/gelatin and Alg-DA/Ac-β-CD/gelatin hydrogels were monitored at 37°C for 24 h. As shown in Fig. 2 a, all groups of hydrogels showed gradually increased swelling ratios within 2 h of incubation, and then the swelling ratios tended to be stable. Compared to unmodified group of hydrogels, all Alg-DA/Ac-β-CD/gelatin hydrogels showed decreased swelling ratio. And the Alg-DA-3/Ac-β-CD/gelatin hydrogel showed a lower swelling ratio of 9.62, which indicated that the swelling ratio of the hydrogel decreased because of the dopamine units
in the hydrogel system. Besides, as shown in Fig. S1 c, the volume of all groups of hydrogels slightly increased during the incubation, thus, the hydrogel could be implanted into defect area without obvious variation of volume.

3.3. In vitro degradation test

The implanted hydrogels are usually required to degrade properly for new tissue ingrowth, achieving a better regeneration of defect tissue. Thus, degradability is an important factor for the implanted hydrogels. In this study, biodegradation behaviors of the prepared hydrogels were measured in PBS and Type-I collagenase/PBS (0.1 mg/mL) solution at 37 °C for 5 days. Degradation test of the samples were conducted by measuring the weight loss during the incubation using a precision analytical balance. As shown in Fig. 2 b, the Alg/Ac-β-CD/gelatin hydrogel group incubated in PBS exhibited an approximate 3.62% of degradation rate after 1 d of incubation and differences between any two groups are not obvious. After 5 days of incubation, Alg-DA-3/Ac-β-CD/ gelatin hydrogel showed a relative lower degradation rate. For the Alg/ Ac-β-CD/gelatin hydrogel group incubated in Type-I collagenase/PBS (Fig. 2 c), 41.87% of the samples degraded after 1 d of incubation. With the incubation time increasing, the degradation rate gradually increased. And with the increasing of dopamine substitution ratio, the degradation rates of the hydrogels were reduced. After 5 days of incubation in Type-I collagenase/PBS, the degradation rate of Alg-DA-3/Ac-β-CD/gelatin hydrogel was 87.18%, which was the lowest among others. For the Alg-DA/Ac-β-CD/gelatin hydrogels, the relative low degradation rates might be because of the induced interactions of catechol groups in the Alg-DA.

3.4. Adhesion property and mechanical property

The adhesive tests were evaluated by placing the hydrogel between two glass slides based on the lap-shear stress test. As shown in Fig. 2 d, after the dopamine substitution, the Alg-DA/Ac-β-CD/gelatin hydrogels with different catechol grafting ratios displayed improved adhesion properties compared with the Alg/Ac-β-CD/gelatin hydrogels. Results showed that Alg-DA-3/Ac-β-CD/gelatin hydrogels showed the adhesion strength of 0.631 ± 0.044 kPa, which was significantly higher than other groups of hydrogels. The compression test results (Fig. 2 e) showed that the elasticity by compression was also improved by dopamine substitution. Alg-DA-3/Ac-β-CD/gelatin hydrogels showed Young’s compression modulus of 26.478 ± 2.846 kPa, which was significantly higher than other groups of hydrogels.

3.5. Effects of PEMFs on hBMSCs chondrogenic and hypertrophic gene expression in vitro

Gene expression level of chondrogenic markers SOX9, Col2a1, ACAN and hypertrophic markers MMP13, Col10a1, RUNX2 (Fig. 3 a) was detected by qPCR after chondrogenic differentiation for 7, 14, and 21 days in hydrogels. Our results showed that PEMF significantly upregulated the expression of chondrogenic mRNA, and downregulated the expression level of RUNX2.

Safranin O staining (Fig. 3 b) and immunohistochemistry (Fig. 3 c, d, e) staining demonstrated that PEMF promoted chondrogenic ECM formation starting from day 7. Significantly larger amounts of chondrogenic matrix were found in the PEMF group compared to control group. IHC staining of MMP13 and Col X (Fig. 3 d, e) showed PEMF treatment significantly inhibited formation of MMP13 and Col X. Staining results also demonstrated that hydrogel showed excellent properties for matrix deposition and cells aggregation. Newly synthesized matrix started to penetrate inside the hydrogels from day 7 and large scale of cell aggregation was observed inside the hydrogels starting from day 14.

3.6. Effects of PEMFs on hMSCs chondrogenic and hypertrophic gene expression and ECM formation in vivo

After nude mice transplantation for 14 days, gross view of recovered hydrogels indicated that our hydrogels were highly deformable (Fig. 4 a). Volume of hydrogels in both groups was decreased compared to those before implantation (Average 31.9 ± 5.46 μL in control group, 30.4 ± 2.72 μL in PEMF group, initial volume before implantation: 50 μL), indicating that our hydrogels slowly degraded after implanted in vivo (Fig. 4 c). However, weight and density of the hydrogels were both increased after implantation, indicating that our hydrogels facilitated the large scale of ECM formation and deposition during implantation in vivo. Significantly higher weight and density of hydrogels were found in PEMF groups, indicating that PEMF treatment promoted ECM deposition. In comparison between control and PEMF groups, mechanical test demonstrated that significantly higher Young’s modulus and ultimate strength were observed in PEMF group, likely resulted by the enhanced ECM formation from PEMF treatment (Fig. 4 d).

qPCR and staining were performed to determine mRNA expression level (Fig. 4 e) and ECM formation (Fig. 4 b, f, g and h) of the hydrogels. In consistent with in vitro results, qPCR and staining results showed that PEMF significantly promoted the gene expression and protein formation of chondrogenic markers and downregulated the expression of hypertrophic markers. Results indicate that PEMF treatment could provide chondro-inductive effects as well as chondroprotective potential during chondrogenesis.

3.7. Bioinformatic analysis of RNA-sequencing and validation study

To better understand the biological mechanisms associated PEMF treatments, hBMSCs from three patients were subject to RNA-sequencing with or without 7 days of PEMF exposure. 17388 transcripts were detected and analyzed, and among which, 54 DEGs (Fold change>2, P < 0.05) were identified. To further investigate the mechanisms dictating the effects of PEMF on BMSCs, KEGG pathway enrichment analysis was performed. The top KEGG pathways were selected based on the most significant fold enrichment (Fig. 5 a, b). The gene-pathway interactions were further visualized in Cerebral Layout using ClueGo (Fig. 5 c). From the bioinformatic analysis, and in combination with the effects observed in vitro and in vivo, we found TNF-α signaling pathway as a potential target pathway in PEMF treatment.

To investigate TNF-α signaling pathway was involved in the effects of PEMF treatment, we visualized the gene expression changes of cells from three individual donors in KEGG pathview [41, 42] (Fig. 5 d). From there, we identified that the two most important MAP kinases, extracellular signal-regulated protein kinase (ERK) and p38 kinase, were most likely to be the key factor of PEMF effects. Further validation using Western blot was performed (Fig. 5 e, f). Results showed that after chondrogenic induction without PEMF treatment, activation and phosphorylation of ERK1/2 were observed on day 1 and peaked on day 3. The expression level of p-ERK1/2 decreased on day 7 and 10. Expression level of p-p38 started low at day 1, kept increasing and reached highest level on day 7 and 10. After PEMF treatment, phosphorylation of ERK1/2 and p38 displayed a similar pattern as control group with continuous promoting of p-p38 and inhibition of p-ERK1/2.

To further validate the involvement of ERK and p38-MAPK pathways, pharmacological inhibition of ERK and p38 was given to hBMSCs pellets during chondrogenic induction and PEMF treatments. qPCR was performed to investigate the expression level of chondrogenic and hypertrophic markers after ERK and p38 inhibition at both early (day 3) and late (day 14) stage of MSCs chondrogenic differentiation (Fig. 5 g).

In control group on day 3, inhibition of ERK1/2 led to increasing expression of Sox9 and RUNX2, and inhibition of p38 resulted in downregulation of Sox9 expression and upregulation of RunX2. On day 14, expression of Sox9 was increased by ERK1/2 inhibition, and decreased by p38 inhibition, and expression of RUNX2 was decreased by
Fig. 3. PEMF affected RNA expression and matrix formation of rBMSCs during chondrogenic differentiation in hydrogels in vitro. a. qPCR analysis results showed that compared to control group, PEMF significantly upregulated the expressions of chondrogenic mRNA, and downregulated the expressions of hypertrophic mRNA. Staining of Safranin O (b) and immunohistochemistry of Col II (c, e) demonstrated that after PEMF treatment, significantly larger amounts of aggrecan and Col II were deposited in hydrogels. Immunohistochemistry of Col X and MMP13 (d, e) demonstrated that PEMF treatment could significantly inhibit the formation of hypertrophic proteins, revealing its chondro-protective potential. Scale bar: 50 μm *p < 0.05 **p < 0.01.
ERK1/2 inhibition, and increased by p38 inhibition. In PEMF group, inhibition of p38 resulted in dramatic downregulation of Sox9 expression on both day 3 and 14, which showed no significant difference compared with control group, indicating that PEMF treatment promoted chondrogenesis of hBMSCs via upregulating the expression and phosphorylation of p38 and mediating p38 MAPK signaling pathway. Inhibition of ERK1/2 resulted in upregulation on day 3 and downregulation on day 14 of RUNX2 expression, which showed no significant difference compared with control group treated with SB203580, indicating that ERK pathway might be involved in the inhibiting effects of PEMF on hypertrophic markers.

3.8. Healing outcomes of cartilage defects

H&E, safranin O and IHC staining (Fig. 6) were performed after 8 weeks of surgery. H&E results demonstrated that most of the hydrogels were dispersed and mostly degraded (Fig. 6 b). The residual hydrogels were surrounded by bony tissues under the newly formed cartilage. Samples from blank and MSCs groups only contained a layer of fibrous tissue with large amount of collagen type I but few collagen type II formation on the surface of defect site (Fig. 6 d, e). In the P-MSCs group, larger amounts of fibrocartilage-like tissue were observed on the surface of defect site. IHC staining demonstrated that the deposited matrix contained both collagen type I and II, confirming the regenerated cartilage to be mostly fibrocartilage. P-MSCs + PEMF group showed the best cartilage healing outcome with thicker layer of hyaline-like cartilage on the defect site. Hyaline chondrocytes were observed in regenerated cartilage surface according to histological staining, and the regenerated matrix on the surface was mostly collagen type II. Furthermore, IHC staining of collagen type X (Fig. 6 f, j) showed that much less collagen type X formation on the defect site were detected in P-MSCs + PEMF group compared to other groups, indicating that PEMF in vivo treatment provides chondroprotective effects against chondrocyte hypertrophy and cartilage degeneration. The quality of the cartilage regeneration is evaluated according to Wakitani scoring system [40]. P-MSCs + PEMF group achieved lowest score, demonstrating the best healing outcomes of osteochondral defect. Rats from P-MSCs group were scored significantly lower than blank and MSCs groups, indicating that PEMF enhanced engineered constructs are more beneficial for cartilage and subchondral bone repair. P-MSCs + PEMF group achieved better healing outcomes than P-MSCs group, indicating that PEMF in vivo treatment also enhanced cartilage regeneration.

4. Discussion

In this study, Alg-DA/β-CD/gelatin hydrogels based on chemical and physical multiple crosslinking and interactions strategy were successfully prepared. After confirming the effect of PEMF signals on hBMSCs chondrogenesis in our hydrogel system both in vitro and in vivo, we demonstrated the therapeutic potential of PEMF on enhancing
cartilage repair in combination of MSCs-based tissue engineering using a rat osteochondral defect model.

Suitable scaffolds are essential for cartilage tissue engineering. In our study, we successfully designed a convenient approach to produce supramolecular hydrogels of desirable attributes via chemical and physical multiple crosslinking and interactions strategy, and thereby introduced stronger and more suitable physical properties for hydrogels in cartilage repair. Our study demonstrated that our hydrogels presented injectable, re-moldable and ideal mechanical properties. Furthermore, owing to the enhanced physical properties brought by chemical and physical multiple crosslinking and interactions, our hydrogels showed relative low swelling ratio, appropriate degradation rate, and tissue adhesive abilities, and have an impact on the availability of implanted hydrogels and benefit their biomedical applications.

3,4-dihydroxyphenethylamine, also called dopamine (DA), which contains catechol group, is a neuromodulatory molecule and exhibit favorable biocompatibility and high adhesive property [43]. DA was often utilized in tissue engineering researches because it showed ideal interactions with tissues via hydrogen bonds and covalent bonds [44]. Therefore, introduction of DA would introduce multiple interactions to the hydrogel system and help to keep the implanted hydrogel stable in the defected tissues.

In our hydrogel system, reversible linkage of the host-guest interactions and chemically crosslinked acrylated structure were the main
structure. With the modification of dopamine, hydrogen bonds and π-π interactions would be introduced into the hydrogel system. But the introduced interactions would not affect the polymerization of double bonds in Ac-β-CD. The aromatic rings of catechol groups could exhibit absorption under 280 nm. But in our hydrogel system, 365 nm of UV light was utilized to initiate the polymerization. Thus, the influence of catechol groups would not be obvious.

It is worth noting that during hydrogel preparation, there is a pre-gel state where increased viscosity was observed. In this state, the pre-gel is in semi-solid form and can be either administrated in situ and then crosslinked, or pre-formed in vitro and injected to the target site, which brought out more translational value to our hydrogel system. This feature grants multiple administration pathways to our hydrogel system, which makes it more practical and can be involved in much more complex biomedical application scenarios. Furthermore, our study demonstrated that the modified hydrogels revealed suitable biological properties such as supporting encapsulated cells, allowing molecular transport and facilitating matrix deposition. Large scale of cell aggregation was observed in our hydrogels in vitro and in vivo, indicating the reversible nature of our hydrogels supports cell migration, which benefited chondrogenesis of MSCs. After transplanted into rat osteochondral defect site, the desirable properties of our hydrogels led to desirable outcomes of subchondral bone healing and cartilage-bone junction formation.

Adult articular cartilage exhibits little capacity for intrinsic repair, and cartilage tissue engineering remains the most promising strategy for the successful repair of articular cartilage defects. By manipulating cells, scaffolds, and stimuli, tissue engineering promised healing of damaged tissues and organs via the use of living, functional constructs [45]. Cartilage, with homogenous structure and few cell types, was originally recognized as one of the first targets of tissue engineering [46]. However, the complex requirement for cartilage regeneration led the healing of cartilage defects to be elusive. In MSCs-based tissue engineering strategies, the chondro-differentiation ability of MSCs was predicted to fulfill the desired criteria for cartilage regeneration. However, apart from chondrocytes, MSCs can also differentiate into other cell types including fibrochondrocytes and hypertrophic chondrocytes, resulting in a mixture of cartilaginous, fibrous, and hypertrophic tissues [19,20]. In vitro chondro-induction of MSCs also results in an unnatural differentiation with co-expression of hyaline cartilage markers (collagen type II and SOX-9) and hypertrophic markers (collagen type X and MMP13) [47]. The success of MSC-based techniques may remain limited if the presence of fibrous and hypertrophic tissue cannot be eliminated [46]. Therefore, in our study, PEMF treatment revealed great therapeutic potential, as it could not only promote chondro-differentiation of MSCs, but also inhibit the expression of hypertrophic markers during chondro-induction, thus enhancing the quality of tissue engineered constructs. Applying PEMF in vivo could also mitigate the hypertrophic process, and facilitate the in situ regeneration of hyaline-like cartilage in osteochondral defects. The effects of PEMF on chondrocytes and MSCs chondrogenic differentiation have been studied, and most of the studies reported the positive effects. The beneficial effects of PEMF on cartilage cells include promoting proliferation, increasing anabolic activities, and antagonizing the catabolic effects of inflammation [22–24]. PEMF treatment affected anabolic activities through upregulating the expression of chondrogenic markers and increasing synthesis of cartilage extracellular matrix (ECM) components, such as GAG and type II collagen [48]. PEMF exposure decreased the pro-inflammatory cytokine release such as interleukin (IL)-6 and IL-8 [25]. Several studies have confirmed the promoting effects of PEMF on MSCs chondrogenic differentiation [49–52]. When exposed to PEMF, type II collagen and glycosaminoglycan (GAG) synthesis were increased in MSCs, indicating that PEMF was able to stimulate and maintain chondrogenesis of MSCs [51]. PEMF exposure during chondrogenic differentiation inhibited the catabolic effects induced by IL-1β, suggesting that PEMF may help to improve clinical outcome in cartilage tissue engineering [53]. More recently, Stafani et al. demonstrated that PEMF exposure could enhance the quality of tissue-engineered cartilage grafts, and therefore promote
engineered cartilage growth and repair [54]. Despite the promising potentials of PEMF that have been reported in chondrogenesis and cartilage repair, the effects of PEMF were believed to be signal specific [55]. Therefore, experimental and clinical studies are still needed to understand the optimal PEMF signal and mechanisms for a defined specific condition. In our study, we used a specific PEMF signal which revealed superior beneficial effects on BMSCs chondrogenesis and cartilage repair. PEMF treatment via this signal promoted rBMSCs chondrogenesis and slowed down cell hypertrophy, indicating its potential to enhance the quality of engineered construct during in vitro preparation. In rat osteochondral defect models, PEMF treatment demonstrated promising therapeutic potential on cartilage repair, and combining with chondrogenic induced MSCs-laden hydrogels, better outcome of cartilage healing was achieved. Through bioinformatic analysis and the following mechanism study, we also demonstrated that our PEMF signal promoted chondrogenesis via upregulating the expression and phosphorylation of p38, and inhibiting hypertrophic markers expression via ERK pathway. These findings suggested that our PEMF signal revealed superior potential in cartilage repair, providing a potential low-cost, low-risk, noninvasive treatment modality to augment the therapeutic effects of cartilage tissue engineering.

In musculoskeletal system, tissues and cells were found sensitive to biophysical stimulations. The diamond concept [21] was thereby proposed to underpin tissue engineering, which introduced environmental biophysical stimulation as the 4th key element apart from cells, scaffolds, growth factor/cytokines. In comparison to other biophysical stimulations, such as mechanical loading and low intensity pulsed ultrasound (LIPUS), PEMF offers unique advantages as a non-contact, non-invasive stimulation. Furthermore, compared to optical, acoustic, and electrical fields as well as mechanical forces, electromagnetic fields have advantages because of their large force output, high precision, and deep tissue penetration [56]. During tissue engineered constructs preparation in vitro, PEMF treatment offers chondro-inductive effects without any increased risk of contamination. As for clinical treatments, since PEMF have been successfully employed as adjunctive therapy for the treatment of many orthopaedics conditions [57–59], numerous types of portable devices were developed, which is suitable for treatment in hospital and physiotherapy from home. This makes PEMF safer and more convenient to use compared to other physical treatments, which is more practical in clinical situations. We therefore consider PEMF to be a perfect candidate as physical stimulation to complete the diamond concept.

Despite our encouraging findings which could lead credence to clinical application, there are limitations in the current study. Firstly, daily PEMF treatment (3 h/day) was given according to our previous studies [32,60]. The dose–response of PEMF treatment in vitro and in vivo was not investigated. Secondly, the parameters of PEMF signal were not investigated in this study, comparison of different PEMF signals might help us further explore more therapeutic signals and understand how PEMFs work and their underlying mechanisms. Future research endeavors should focus on these studies, in order to explore more practical tissue engineering strategies for clinical applications.

5. Conclusion

The chemical structure, injectability, self-healing behavior, morphology, swelling behavior, adhesiveness, in vitro/in vivo biocompatibility, and therapeutic efficacy for cartilage repair of the Alg-DA/Ac-β-CD/gelatin hydrogels were systematically characterized. All of the results suggested that the injectable self-healing adhesive hydrogel exhibited ideal properties for cartilage engineering. PEMF signal promoted BMSCs chondrogenesis and inhibited hypertrophic processes, and thereby enhanced the quality of engineered chondrogenic constructs. Together with enhanced chondrogenic constructs induced from BMSCs laden hydrogels, PEMF promoted cartilage healing in rat osteochondral defect models, and demonstrated great potential as 4th element in cartilage tissue engineering and translational value in clinical applications.

CRediT authorship contribution statement

Yucong Li: Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Project administration, Roles/Writing – original draft. Linlong Li: Data curation, Formal analysis, Investigation, Methodology, Roles/Writing – original draft. Ye Li: Data curation, Formal analysis, Investigation, Roles/Writing – original draft. Lu Feng: Investigation, Resources, Software. Bin Wang: Investigation, Resources, Software. Ming Wang: Investigation, Methodology. Haixing Wang: Investigation, Methodology. Meiling Zhu: Methodology. Yongkang Yang: Investigation. Erik I. Waldorff: Conceptualization, Funding acquisition. Nianli Zhang: Conceptualization, Funding acquisition. Ingmar Viohl: Resources, Funding acquisition. Sien Lin: Supervision, Validation, Writing – review & editing. Liming Bian: Supervision, Validation. Wayne Yuk-Wai Lee: Supervision, Validation, Writing – review & editing. Găng Li: Conceptualization, Funding acquisition, Project administration, Resources, Writing – review & editing.

Declaration of competing interest

The authors have no conflicts of interest to disclose in relation to this article. NZ, EIW and IV are employees of and own stock in Orthofix Medical Inc., USA.

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Appendix A. Supplementary data

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