Research Article

Multifaceted Optimization of MSC-Based Formulation upon Sodium Iodoacetate-Induced Osteoarthritis Models by Combining Advantageous HA/PG Hydrogel and Fluorescent Tracer

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Owing to the boundedness of conventional remedies upon articular cartilage for self-rehabilitation and the incrementally senior citizens, the incidence of osteoarthritis (OA) is increasing worldwide. Empirical studies have revealed the advantageous and promising potentials of mesenchymal stem/stromal cells (MSCs) on the refractory OA, whereas the deficiency of systematic and detailed exploration of MSC-based therapy largely hampers the large-scale applications in regenerative medicine. Herein, we initially utilized the monosodium iodoacetate- (MIA-) induced OA rabbit models and investigated the therapeutic effect of human umbilical cord-derived UC-MSCs at serial dose gradients with the splendid hyaluronic acid and/or propylene glycol hydrogels (HA, HA/PG), respectively. Afterwards, we turned to a dual-luciferase reporter tracing system and evaluated the spatiotemporal distribution and metabolokinetics of bifluorescence expressing UC-MSCs (BF-MSCs) in OA rats. Of the aforementioned trials, we verified that the combination of HA/PG and middle-dose MSCs (0.5 × 10^7 cells/ml) eventually manifested the optimal efficacy on OA rabbits. Furthermore, with the aid of the bioluminescence imaging (BLI) technology for dynamic in vitro and in vivo tracking, we intuitively delineated the spatiotemporal distribution and therapeutic process of BF-MSCs in OA rats, which substantially confirmed the reinforcement of HA/PG on BF-MSCs for OA treatment. Collectively, our data conformably demonstrated that the middle dose of UC-MSCs combined with HA/PG hydrogel was sufficient for optimal MSC-based formulation for blocking OA progression and promoting cartilage repair, which supplied overwhelming new references and enlightened MSC-based therapeutic strategies for cartilage defects.

1. Introduction

Osteoarthritis (OA) is acknowledged as the most prevalent and frequent chronic disease of degenerative joints that attributes to multidimensional factors such as aging, over-weight, trauma, congenital abnormality, and joint deformity and, in particular, the increase in life expectancy [1, 2]. Despite the dramatic progress and tremendous effects have been made by multidisciplinary strategies, including surgeries (e.g., microfracture and mosaicplasty), anti-inflammatory drugs (e.g., NSAIDs), joint lubricants (e.g., hyaluronic acid), and even the cell-based therapies (e.g., autologous chondrocyte implantation), yet the limitation of self-repair and regeneration capacity in OA individuals still hinders the remission of the progressive degeneration of articular cartilage [3–5].
Mesenchymal stem/stromal cells (MSCs) are heterogeneous and immune-privileged populations with multidirectional differentiation potential (e.g., adipocytes, osteoblasts, chondrocytes, and angiogenic), which are therewith recognized as the uppermost components in the hematopoietic microenvironment as well as advanced sources for translational medicine [6, 7]. For decades, we and other investigators have illuminated the identification and application of the advantageous and promising sources from adult and perinatal tissues together with human pluripotent stem cells (hPSCs) for the treatment of refractory and recurrent disorders such as aplastic anemia, acute-on-chronic liver failure (ACLF), fistulizing Crohn’s disease, acute myocardial infarction (AMI), critical limb ischemia (CLI), and coronavirus disease 2019- (COVID-19) induced acute respiratory distress syndrome (ARDS) [8–13]. State-of-the-art updates have indicated the possibility of MSCs as an alternative option for cartilage repair, especially the allogeneic UC-MSCs with advantageous superiorities in proliferation and immunomodulation and noninvasiveness [12, 14]. Although administration of OA in preclinical trials by utilizing UC-MSCs has been partially achieved, yet the resumption of the functionally competent articular cartilage is far from satisfaction, let alone the meticulous evaluation of MSCs upon safety and efficacy, which is also the prerequisite for large-scale application in OA treatment [4, 15].

For the purpose, we incipiently took advantage of the well-established MIA-induced OA rabbit model and verified the optimally curative effect by comparing the combination schemes among HA/PG and/or UC-MSCs at various gradients (low, middle, or high dose). With the aid of systematic and detailed assessments including clinicopathological observation of articular defects and disease score of multiple gradients (low, middle, or high dose). With the aid of systematic and detailed assessments including clinicopathological observation of articular defects and disease score of multiple

2. Materials and Methods

2.1. Rabbits. According to Wang et al. and Hartman et al. reports, with the increase of age and the period of osteoarthritis, the biomechanics and metabolic capacity of articular chondrocytes obviously degenerate. Old rabbits are prone to cartilage degeneration, which affects the consistency and pathological changes of the models [16, 17]. Hence, 28 healthy New Zealand white male rabbits (2.0 ± 0.5 kg) were purchased and quarantined for further experimental analyses (YUDA Biotechnology Co., Ltd, Tianjin, China). All procedures were approved by the Ethics Committee of Eye Hospital of Tianjin Medical University (approval number: TJYY2018061114), and the in vivo fluorescence tracer experiments were performed with SD rats and approved by the Institutional Animal Care and Use Committee of Institute of Radiation Medicine, Chinese Academy of Medical Sciences & Peking Union Medical College. For each SD rat, anesthesia was conducted with isoflurane; both legs were then injected by using MIA as previously described [18, 19]. The total volume of UC-MSCs (100 μl per joint, 0.5 × 10⁶ cells/ml) was injected into the joint of the knee. Meanwhile, we confirmed that all procedures on rabbits and rats were performed in accordance with the Helsinki Declaration of 1975 (as revised in 2008) concerning Human and Animal Rights.

2.2. MIA-Induced Osteoarthritis Rabbit Models. The MIA-induced osteoarthritis rabbit model was displayed as previously reported with several modifications [15, 20, 21]. Briefly, 28 rabbits were randomly divided into seven groups as the following: (i) negative control group (NC, without any treatment; n = 4), (ii) saline group (MIA-induced OA, with saline injection; n = 4), (iii) HA group (MIA-induced OA, with HA injection; n = 4), (iv) HA/PG group (MIA-induced OA, with HA/PG injection; n = 4), (v) HA/PG+UC-MSC (high) group (MIA-induced OA, with HA/PG and 1.0 × 10⁶ cells/ml injection; n = 4), (vi) HA/PG+UC-MSC (middle) group (MIA-induced OA, with HA/PG and 0.5 × 10⁶ cells/ml injection; n = 4), and (vii) HA/PG+UC-MSC (low) group (MIA-induced OA, with HA/PG and 0.1 × 10⁶ cells/ml injection; n = 4). All applicable institutional and/or national guidelines for the care and use of animals were followed. The detailed procedures and primer sequences were available in Supplementary Information: Supplementary Procedures and Supplementary Table S1.

2.3. Preparation of UC-MSC and HA/PG Hydrogel Composite. UC-MSCs were purchased from the cell products of the National Engineering Center & National Stem Cell Engineering Research Center (Tianjin IMCELL Stem Cell and Gene Technology Co., Ltd., Tianjin). Various concentrations of UC-MSCs at passage three (0.1 × 10⁷ cells/ml, 0.5 × 10⁷ cells/ml, and 1.0 × 10⁷ cells/ml) were thoroughly mixed with 1% sodium hyaluronate/proplylene glycol (2:1, Shandong Freda Biotechnology Co., Ltd, China; HA: Mw = 1579450 Da, with the requirements of Chinese Good Manufacturing Practices for Pharmaceutical Products) for MSC/hydrogel composite generation. The detailed procedures and antibodies for FCM assay were available in Supplementary Information: Supplementary Procedures and Supplementary Table S1-S2.

2.4. Macroscopic Analysis. Macroscopic analysis of femoral condyles and tibial plateau in OA rabbits was conducted according to the criteria as reported [22]. In detail, the aforementioned knee joints were carefully collected after euthanasia for the examination of the severe inflammation or extensive fibrosis in the joint. Then, the degree of femur and tibia condyle repair of each rabbit was macroscopically assessed by means of the modified version of the Osteoarthritis Research Society International (OARSI) scoring system [22]. In detail, 3 independent observers graded the OARSI by utilizing the randomized and blind methods. The consistency of these observers was determined by the intragroup correlation coefficient, which was used for
the assessment of the correlation between the same single pathological section.

2.5. Histopathological Analysis. The preparation for histopathological analysis was displayed as we recently reported with several modifications [6]. Full-thickness samples (including femoral condyles and tibial plateau) were orderly fixed, decalcified, paraffin-embedded, and deparaffinized. For evaluation of OA, microphotographs of the sections were taken using a light microscope (Olympus, BX51, Japan) after hematoxylin and eosin (H&E), Safranin-O/fast green or Alcian blue staining (Beijing Solarbio Science & Technology Co., Ltd. China), respectively. The grade of osteoarthritic change was assessed based on the staining areas according to Mankin’s score and the formula: SEM percentage = (SEM/Mean) × 100 [23]. The detailed procedures and qRT-PCR analysis were available in Supplementary Information: Supplementary Procedures and Supplementary Table S1.

2.6. Cytokine Expression in Synovial Fluid of OA Rabbits. To detect OA-associated inflammatory cytokine and growth factor expression, synovial fluid was collected from each knee joint of rabbits. Firstly, we conducted the preparation of synovial fluid as the following: 20x dilution of washing solution: 1:20 dilution with double distilled water (e.g., 1 ml of concentrated washing solution was added with 19 ml of double-distilled water). Then, the concentrations of interleukin-6 (IL-6), interleukin-1β (IL-1β), transforming growth factor-β1 (TGF-β1), and matrix metalloproteinases-13 (MMP-13) were detected with commercial enzyme-linked immunosorbent assay (ELISA) kits (R&D Systems, Inc., USA) according to the manufacturer’s instructions as we described previously [12, 24]. The detailed information was available in Supplementary Information: Supplementary Table S3.

2.7. Principal Component Analysis (PCA). PCA was used to reveal the internal structure of the present multivariate dataset as a set of coordinates in the corresponding high-dimensional data space as we and other investigators reported [6, 25]. The effects of treatment groups and their correlation with the other characteristics were determined statistically by PCA based on the XLSTAT software (Addinsoft, New York, USA).

2.8. In Vivo Tracking of BF-MSCs (with/without HA/P) in OA Rats. In vivo tracking of BF-MSCs in SD rats was conducted based on the bioluminescence imaging (BLI) signal (p/s/cm2/sr) as we and our collaborators recently reported with several modifications [11, 26]. BF-MSCs with firefly luciferase (Fluc) and green fluorescent protein (GFP) expression were prepared in D-luciferin (33 μl/mg; Cayman Chemical, USA) before intra-articular injection. Then, a total volume of BF-MSCs (100 μl per joint, 0.5 × 10⁶ cells/ml) was injected into each knee joint. Images were acquired 1 h, 48 h, 96 h, and 144 h with an exposure time of 10 min. The detailed procedures were available in Supplementary Information: Supplementary Procedures. All applicable institutional and/or national guidelines for the care and use of animals were followed.

2.9. Statistical Analysis. All statistical analyses were performed as we described before [12, 27, 28]. The experiments were performed in triplicate for three times, and data were shown as mean ± standard deviation (SD). For the purpose of dissecting the difference between multiple parameters (including Anderson-Darling test, D’Agostino-Pearson omnibus normality test, Shapiro-Wilk normality test, and Kolmogorov-Smirnov normality test with Dallal-Wilkinson-Lilliefors P value) of knee joints (8 knee joints in 4 rabbits per group), we primitively checked the Gaussian distribution and found that their respective P value summary was NS, which satisfied the criteria of single-factor analysis of variance. Thus, one-way analysis of variance (ANOVA) was used for statistical analysis. P values were considered significantly when P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001; NS means not significant.

3. Results

3.1. Moderate UC-MSC and HA/P Composite Effectively Ameliorated Pathological Damage of Articular Cartilage in OA Rabbits. For the purpose of systematically and meticulously dissecting the curative effect and optimizing the MSC-based formulation upon osteoarthritis, we primordially took advantage of the well-established sodium iodoacetate-induced rat model [15, 20, 21] (Figure 1(a)). After euthanasia at week 13, the intact femur and tibia of the knee joints in rabbits among the indicated groups were carefully obtained by surgical operations as described in Materials and Methods. Distinguish from those in the saline group (OA rabbit with saline injection), the joint surface of tibia showed partially ameliorated pathological damage of articular cartilage after injection of UC-MSCs or HA/P hydrogel, especially OA rabbits with HA/P+high or HA/P+middle dose UC-MSC administration (Figure 1(b)). Similarly, according to the gross morphological observations upon cartilage abrasion and subchondral bone exposure, MIA-induced OA rabbits with intra-articular injection of HA/P+high and HA/P+middle dose UC-MSCs manifested comparable restorative function in the femur as well (Figure 1(c)).

To further quantitatively determine the ameliorative pathological damage of articular cartilage, we turned to OA scoring based on the OARSI macroscopic evaluation system and confirmed that rabbits with HA/P+high or HA/P+middle dose UC-MSC injection showed significantly lower scores than those in the other groups, whereas indistinguishable macroscopic differences were observed between these two groups [22] (Figure 1(d)). Interestingly, we found the OARSI macroscopic score of femurs in rabbits with HA/P+middle dose UC-MSCs was further decreased compared with the HA/P+high dose UC-MSC group, which was in consistence with smoother morphological characteristics of joints (Figures 1(d) and 1(e)). Collectively, these data suggested that moderate UC-MSC and HA/P composite showed superiority in ameliorating macroscopically pathological damage of articular cartilage in OA rabbits.

3.2. Clinicopathological Assessment of the Osteochondral Defects Confirmed the Preferable Efficacy by Moderate UC-
Establishment of osteoarthritis model and HA/PG/UC-MSC transplantation

2 w 4 w 6 w 8 w 13 w

Saline/HA/PG/UC-MSCs Bilateral i.a injection twice

2% MIA (100 μl) Bilateral i.a injection twice

Macroscopic analysis
Histological analysis
Cytokine analysis
PCA analysis (optimum formula)

(a)

NC  Saline  HA  HA/PG  HA/PG+hUC-MSCs (dose)

Tibia
Left

Right

HA/PG+UC-MSCs (dose)

High  Middle  Low

(b)

NC  Saline  HA  HA/PG  HA/PG+hUC-MSCs (dose)

Femur
Left

Right

(c)

Statistical analysis of the OARSI macroscopic score of tibias (d) and femurs (e) in the aforementioned 7 groups. *P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001; NS: not significant.

Figure 1: Pathological changes of articular cartilage in OA rabbits with UC-MSC and/or HA/PG composite administration. (a) Schedule of the MIA-induced OA rabbit model with UC-MSC and/or HA/PG composite administration. (b, c) Representative macroscopic images of the tibias (b) and femurs (c) in the indicated groups with intra-articular (i.a.) injection of saline, HA, HA/PG, high/middle/low dose of UC-MSCs+HA/PG hydrogels at week 13, respectively. Rabbits without any treatment were served as negative controls (NC). (d, e) Statistical analysis of the OARSI macroscopic score of tibias (d) and femurs (e) in the aforementioned 7 groups. *P < 0.05; **P < 0.01; ***P < 0.0001; ****P < 0.00001; NS: not significant.
MSC and HA/PG Composite. Having preliminarily identified the ameliorative efficacy of moderate UC-MSC and HA/PG composite in macroscopically pathological damage, we subsequently analyzed the potential distinctions among the indicated groups at a histological level. As shown by the representative photomicrographs of femoral condyles and tibial plateau with H&E, Alcian blue, and Safranin-O/fast green staining, rabbits with HA/PG+middle or HA/PG+high dose UC-MSC treatment exhibited less cartilage loss and surface irregularity of the joints as well as much typical and homogenous articular cartilage structures including smooth, continuous surfaces and strong proteoglycan staining (Figures 2(a)–2(c)). In contrast, distinguish from the abovementioned two groups and the NC group, the surface irregularity, fibrillation or cleft, changes in cellularity, and loss of tidemark were observed in varying degrees among the other MIA-induced OA groups (saline, HA, or HA/PG group) (Figures 2(a)–2(c)).

Furthermore, quantitative analyses of the histopathologic sections of articular cartilage in OA rabbits were performed by measurement of the percentages of areas with specific staining as we previously reported and also described in the supplementary information [25]. As shown by the calculations based on the aforementioned H&E staining, HA/PG hydrogel combined with either dose (low, middle, and high)
of UC-MSCs exhibited a significantly higher percentage of stained areas compared with the saline group, whereas similar to that in the NC group (Figure 3(a)). As to Alcian blue staining, rabbits received HA/PG+middle dose of UC-MSC treatment exhibited a significantly higher proportion of positive area for collagen staining when compared with the saline group. However, no significant differences were observed between HA/PG+middle dose of UC-MSC and NC groups (Figure 3(b)). Consistently, we found the tendency of the proportions of positive areas with Safranin-O staining. All data were shown as mean ± SD (n = 8). *P < 0.05; **P < 0.01; ***P < 0.001; NS: not significant.
staining among the indicated groups was similar to that with the Alcian blue staining. That is, rabbits with HA/PG+middle dose of UC-MSC administration exhibited a significantly higher percentage of positive area for proteoglycan-associated GAG staining than the other OA groups (Figure 3(c)). Simultaneously, we aimed to further dissect the potentially microscopic alterations of articular cartilage based on the modified Mankin score as well. In regard to the Mankin score of H&E staining, rabbits with HA/PG hydrogel+UC-MSC injection revealed lower scores than the other groups (saline, HA, or HA/PG) to some extent (Figure 3(d)). Different from those with a high or low dose of UC-MSC treatment, rabbits with middle dose showed further decreases in Mankin scores of Alcian blue staining and Safranin-O staining, respectively (Figures 3(e) and 3(f)).

Taken together, these data further confirmed that rabbits with HA/PG+middle dose of UC-MSC administration exhibited a significant reduction of proinflammatory cytokines (IL-1β, IL-6, MMP-13), which indicated the superiority in anti-inflammatory property (Figure 4(d)). Collectively, these data manifested enhanced residence of UC-MSCs upon OA.

3.4. Establishment of Bifluorescence Expressing BF-MSCs with Signatures of Immunophenotype and Multilineage Differentiation. For spatiotemporal dissection of the metaboolokinetics of UC-MSCs on OA treatment, we utilized the bioluminescence imaging (BLI) technology with the lentivirus-mediated dual-fluorescence reporter construct (pLV-Fluc-eGFP) as we and collaborators reported recently [11, 26, 29, 30] (Figure 5(a)). After continuous enrichments, UC-MSCs expressing a high level of luci and GFP were established (denoted as BF-MSCs), which also exhibited representative immunophenotype of adult MSCs (Figures 5(b) and 5(c)). Concurrently, by conducting multilineage differentiation detection, we found the BF-MSCs were sufficient to, respectively, generate adipocytes, osteoblasts, and chondrocytes under the indicated conditions as shown by the O Red O, Alizarin Red, and Alcian blue staining and qRT-PCR analysis of adipogenic (PPARγ and ADIPOQ), osteogenic (RUNX2 and BGLAP), and chondrogenic (ACAN and SOX9) associated genes (Figures 5(d) and 5(e)). In addition, with the aid of amplification curve and karyotypic analysis, we verified that the BF-MSCs after gene editing exhibited admirable in vitro proliferation and chromosomal stability as well (Figures 5(f) and 5(g)). Taken together, the established BF-MSCs with bifluorescence expression possessed decent signatures for further in vivo analysis.

3.5. BF-MSCs Exhibited Stable Fluorescence Intensity Both In Vitro and In Vivo. To further evaluate the sensitivity of the GFP and Fluc fluorescences in BF-MSCs as well as the potentiation effect of the HA/PG hydrogel, we incipiently performed an in vitro dosage-dependent experiment by seeding cells into 12-well plates at various density gradients. In consistence with our previous reports, both the GFP and BLI signal intensity of Fluc showed pertinently matched intensity with the density of BF-MSCs in vitro under pretreatment with firefly luciferase substrate D-luciferin (Figures 6(a)–6(c)) [8, 29].

For the purpose of verifying the spatiotemporal metabolokinetics of BF-MSCs as well as the enhanced therapeutic effect of the moderate UC-MSC and HA/PG composite in vivo, we turned to another well-established MIA-induced OA model on SD rats (Figure 6(d)). Different from those in the control group with only UC-MSC injection (denoted as -HA/PG), rats with moderate UC-MSC and HA/PG composite administration showed delayed decline in BLI signal according to the intensity gradients of BLI signal by in vivo imaging systems (IVIS), which suggested the credibility of potentiation effect of HA/PG to BF-MSCs. Additionally, the transplanted UC-MSC and HA/PG composite could maintain in the sites of articular cartilage in OA rats and survive for one week (Figures 6(d) and 6(e)). Therefore, the BF-MSCs was also sensitive enough for spatiotemporal distribution and metabolism analysis of cell tracking in vivo, and the HA/PG hydrogel manifested enhanced residence of UC-MSCs upon OA.

4. Discussion

Osteoarthritis is a principal and challenging disorder of articular cartilage due to multiple pathogenic factors and a broad
Figure 4: Expressions of proinflammatory cytokines in joint fluid and systematic assessment of OA rabbits. (a–d) Statistical analysis of the expression levels of proinflammatory factors in synovial fluid in the indicated 7 groups by ELISA including IL-1β (a), IL-6 (b), MMP-13 (c), and TGF-1β (d). Data were shown as mean ± SD (n = 8). * P < 0.05; ** P < 0.01; *** P < 0.001; NS: not significant. (e) Principal component analysis (PCA) of the aforementioned 19 parameters involved in the MIA-induced OA rabbit model among the indicated 7 groups. F1 and F2 represented principal components 1 and 2 (PC-1 and PC-2), respectively.
demographic population, and in particular, the elders are confronting and struggling with the burden of the refractory chronic disease [2, 4, 5]. Even though multifaceted improvements have been achieved by multidisciplinary therapies such as surgical procedures and anti-inflammatory drugs, yet the pathogenesis and efficacy of OA with degeneration and functional losses are still elusively complicated for ascertainment [3, 31]. Herein, we took advantage of the promising MSC-based cytotherapy and advantageous HA/PG hydrogel composites for optimizing the formulation upon MIA-

![Figure 5: The multifaceted characterizations of the dual-fluorescence expressing BF-MSCs.](image)

(a) Illustration of the lentivirus-mediated dual-fluorescence reporter construct (pLV-Fluc-eGFP). Fluc and GFP were for BLI and green fluorescent protein imaging, respectively. (b) The phase contrast and GFP fluorescence of dual-fluorescence expressing UC-MSCs (denoted as BF-MSCs). (c) FCM analysis of GFP and MSC-associated biomarker in BF-MSCs. (d) Multilineage differentiation potential (adipogenic, osteogenic, and chondrogenic differentiation) of BF-MSC-derived cells (upper panel) and untreated DF-MSCs (bottom panel) identified by Oil Red O (left panel), Alizarin red (middle panel), and Alcian blue (right panel) staining, respectively. Scale bar = 30 μm. (e) qRT-PCR analysis of the adipogenic- (ADIPOQ and PPARγ), osteogenic- (RUNX2 and BGLAP), and chondrogenic- (ACAN and SOX9) associated genes in the indicated BF-MSCs (mean ± SD, n = 3). **P < 0.01; ***P < 0.001. (f) Amplification curve of BF-MSCs based on the cumulative cell numbers and days of cultivation (mean ± SD, n = 3). (g) Karyotypic analysis of UC-MSCs and BF-MSCs cultured in 10%FBS/DF12 medium with G-banded chromosome detection.
Fluorescent images of BF-MSCs in a dosage-dependent manner (GFP)

- 6.25 x 10^3
- 1.25 x 10^4
- 2.5 x 10^4
- 5 x 10^4
- 1 x 10^5
- 2 x 10^5

(a)

(b) Luminescence

(c) Radiance (p/sec/cm^2/sr) × 10^6

(d) Correlation of Fluc with cell counts

Figure 6: Continued.
induced osteoarthritis models upon rabbits and rats. Furthermore, aided with the BLI signal, we intuitively and dynamically figured out the spatiotemporal metabolokinetics of MSCs in vivo, which would congruously benefit the fundamental and clinical researches upon the safety and effectiveness assessment of MSC-based cytotherapy.

Of the reported MSCs, UC-MSC is regarded as a preferable source with superiorities including long-term proliferation and immunoregulatory attributes for numerous recurrent and refractory disease remodeling including cartilage regeneration [12, 32, 33]. Meanwhile, we and other investigators recently also reported the discrepant and controversial efficacy of UC-MSCs on disorders such as GVHD, acute liver failure, and MIA-induced osteoarthritis attribute to cell vitality and dosage, respectively [34–38]. Considering the contingent risks in safety and effectiveness, it is of importance and indispensability for systematically and meticulously dissecting the optimal formulation for MSC-based therapy before large-scale clinical applications. Herein, by conducting intra-articular injection of UC-MSCs at various concentration gradients, we found moderate MSC application manifested consistent efficacy but preferable immunoregulatory capacity in suppressing proinflammatory factors in joint fluid. Additionally, the BLI signal of dual-fluorescence expressing BF-MSCs was sufficient for in vivo real-time assessment of cell metabolokinetics.

State-of-the-art progress has highlighted the feasibility and cooperativity of biomaterials with MSCs in regenerative medicine [39, 40]. For instance, Park et al. reported UCB-MSCs and HA hydrogel in repairing articular cartilage lesion in surgery-induced OA rabbit and rat models, respectively [37, 40]. However, by utilizing a more realistic and intractable MIA-induced model with characteristics including proteoglycan matrix loss and disruption of glycolysis as well as functional joint impairment, van Buul et al. found MSC administration could only partially reduce pain without regaining the beneficial effect on degenerative changes [38]. To overcome the shortcomings, we repeated the MIA-induced OA model both on rabbits and on rats and conducted multifaced optimization of MSC-based formulation with the much more advantageous HA/PG hydrogel, which exhibited enhanced therapeutic efficacy than HA and approved as a safe and food-grade ingredient by the Food and Drug Administration (FDA) in ameliorating osteoarthritis progression [41, 42]. Hence, we have conducted a novel composite by moderate UC-MSCs and HA/PG hydrogel to dissect the therapeutic potential in MIA-induced OA models.

Effective evaluations upon safety, effectiveness, and reproducibility in preclinical and clinical studies are the prerequisites for large-scale application of MSC-based therapy, especially the pharmacokinetic and pharmacodynamic characterization in vivo [14, 34]. However, most of the current methodologies for cell labeling including radioprobes and fluorophores were insufficient for long-term cell monitoring due to the excessive dilution of agents during cellular proliferation and persistence of signals from released agents and dead cells [43, 44]. Conversely, we and other investigators have recently developed an indirect and dual-fluorescent labeling system for in vivo MSC tracking based on the lentivirus-mediated GFP and luciferase (Luc), which is competent for tracer upon exosome vesicle as well [11, 26, 45]. In this study, our data showed that the system was also capable of tracing the spatiotemporal metabolokinetics of UC-MSC+HA/PG composite. Above all, MSCs in HA/PG displayed preferable homing and delayed attenuation in the damaged articular cartilage compared with single HA/PG or UC-MSC administration attribute to the splendid viscoelasticity, minimal influence to cell behavior, and the encapsulated microenvironment by HA/PG [8, 11, 41, 46].

5. Conclusions

Overall, our data suggested that moderate UC-MSCs and HA/PG composite supplied an optimal formulation upon
MIA-induced osteoarthritis models and the potential in regenerative medicine for alleviating articular cartilage defects. Meanwhile, the BLI-based system was sufficient for monitoring the dynamic metabolism of UC-MSCs (with/without HA/PG) in vivo, which was promising for the assessment of MSC-based cytotherapy before large-scale application.

Data Availability

The data used to support the findings of the study are included in the article. Additional data related to this study are also available from the corresponding author.

Conflicts of Interest

The indicated commercial companies including Health-Biotech (https://www.health-biotech.com/en/intro.html), Amcellgene (https://www.amcellgene.com/en/), and Chase Sun (http://www.chasesun.cn) gave technical support, which we have expressed our thanks in the “Acknowledgments” section.

Authors’ Contributions

Ai-tong Wang and Meng Zhao performed the experiments, collection, and assembly of data; Ying Feng, Honghong Jia, Zongiin Li, Zhibo Han, and Zhongchao Han helped with collection and assembly of data; Leisheng Zhang and Hao Yu contributed to conception and design, data analysis and interpretation, manuscript writing, and final approval of the manuscript. All authors read and approved the final manuscript. Ai-tong Wang and Meng Zhao contributed equally to this work.

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Supplementary Materials

Supplementary Table 1: the primer sequences of indicated genes for quantitative RT-PCR analysis. Supplementary Table 2: the antibodies for flow cytometry analysis. Supplementary Table 3: the ELISA kits used in this study. Supplementary Procedures: the additional procedures accompanied with the main manuscript. Supplementary References: the additional references accompanied with the main manuscript. (Supplementary Materials)

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