Comparison of Bone Marrow Aspirate Concentrate and Allogenic Human Umbilical Cord Blood Derived Mesenchymal Stem Cell Implantation on Chondral Defect of Knee: Assessment of Clinical and Magnetic Resonance Imaging Outcomes at 2-Year Follow-Up

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Abstract
Biological repair of cartilage lesions remains a significant clinical challenge. A wide variety of methods involving mesenchymal stem cells (MSCs) have been introduced. Because of the limitation of the results, most of the treatment methods have not yet been approved by the Food and Drug Administration (FDA). However, bone marrow aspirate concentrate (BMAC) and human umbilical cord blood derived mesenchymal stem cells (hUCB-MSCs) implantation were approved by Korea FDA. The aim of this study was to evaluate clinical and magnetic resonance imaging (MRI) outcomes after two different types of MSCs implantation in knee osteoarthritis. Fifty-two patients (52 knees) who underwent cartilage repair surgery using the BMAC (25 knees) and hUCB-MSCs (27 knees) were retrospectively evaluated for 2 years after surgery. Clinical outcomes were evaluated according to the score of visual analogue scale (VAS), the International Knee Documentation Committee (IKDC) subjective, and the Knee Injury and Osteoarthritis Outcome Score (KOOS). Cartilage repair was assessed according to the modified Magnetic Resonance Observation of Cartilage Repair Tissue (M-MOCART) score and the International Cartilage Repair Society (ICRS) cartilage repair scoring system. At 2-year follow-up, clinical outcomes including VAS, IKDC, and KOOS significantly improved (\( P < 0.05 \)) in both groups; however, there were no differences between two groups. There was no significant difference in M-MOCART [1-year (\( P = 0.261 \)), 2-year (\( P = 0.351 \))] and ICRS repair score (\( P = 0.655 \)) between two groups. Both groups showed satisfactory clinical and MRI outcomes. Implantation of MSCs from BMAC or hUCB-MSCs is safe and effective for repairing cartilage lesion. However, large cases and a well-controlled prospective design with long-term follow-up studies are needed.

Keywords
cartilage repair, articular cartilage injury, MSC, BMAC, hUCB-MSCs

Introduction
Articular cartilage is known to have low potential for healing, and damage from degeneration or trauma can lead to focal cartilage lesions and subsequent osteoarthritis. Cartilage lesion remains a challenging area in orthopedics, despite various treatment methods having been attempted. The technique of microfracture has shown good short-term clinical results within 2 years; however long-term data for larger and more aged patients have suggested inferior

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established6–8. Therefore, most stem cell treatment methods in studies in terms of treatment methods, number of cells, incubation methods, and paucity of high-quality controlled growth factors, and cytokines6,9,13. These components can cell counts and ability of differentiation, BMAC contains not may not be obtained. Despite limitations in terms of stem

As the cell-based tissue engineering techniques have improved, various cartilage treatment methods using stem cells [including mesenchymal stem cells (MSCs)] have been proposed5,6. Nevertheless, because of a lack of consistency in studies in terms of treatment methods, number of cells, incubation methods, and paucity of high-quality controlled study, no clear standardized treatment methods have been established6–8. Therefore, most stem cell treatment methods (e.g., adipose cell-derived MSCs or adipose cell sourcederived stromal vascular fraction cells) have not yet been approved by the US Food and Drug Administration (FDA). However, bone marrow aspirate concentrate (BMAC) can be prepared in a standardized manner with FDA and Korea FDA (KFDA) approval. Human umbilical cord blood derived mesenchymal stem cells (hUCB-MSCs) have been developed as a drug product (Cartistem®, Medipost Inc., Sungnam, Gyeonggi-do, South Korea) with the approval of KFDA and can be used in Korea, and a clinical trial is underway in the United States6.

BMAC, containing MSCs, used with hyaluronic acid based scaffold has generated satisfactory clinical outcomes and successful cartilage repair9,10. It is fairly easy to perform bone marrow aspiration from the iliac crest through mini-incision without position change. Subsequently, BMAC is generated using commercialized machines. This method can overcome the limitations of cell-based two-stage procedures, promising one-stage option for cartilage repair procedure with chondrogenic potential and easy control of MSCs. Furthermore, the risk of disease transmission is low. Nevertheless, BMAC contains only a small fraction of MSCs. In general, we can extract about 1.9 × 10^7 mononuclear cells of approximately 6 ml capacity after aspirating 60 ml of bone marrow9. The number of collected MSCs may vary depending on the patient’s age and general condition11. MSCs from elderly patients have lower differentiation levels and chondrogenicity in vitro12. For these reasons, satisfactory results may not be obtained. Despite limitations in terms of stem cell counts and ability of differentiation, BMAC contains not only stem cells but also hematopoietic stem cells, platelets, growth factors, and cytokines6,9,13. These components can have substantial paracrine effect14,15.

hUCB-MSCs were first reported by Erices et al16. They have high proliferation rates and can be easily induced to differentiate into chondrocytes. Despite, still under debate on chondrogenic potential of MSC source17–19, hUCB-MSCs have low immunogenicity, and therefore are well suited to allogenic transplantation20. This method has the advantage of low donor site morbidity, no need for preparation procedures, and uniform cell count and quality. Furthermore, it allows implantation of a specific amount of MSCs, regardless of the size of the lesion. Nevertheless, there are risks of contamination or damage because of mistakes in manufacturing or storage prior to use. One study reported satisfactory outcome with 7-year follow-up after hUCB-MSCs transplantation21.

Currently, most reported studies of cartilage repair treatments using the stem cells compared results with those of microfracture10,22. Alternatively, the effects of cartilage regeneration were assessed by comparing the pre- and postoperative status23,24. Because of the limitations of the microfracture technique, it is not appropriate to compare it with the effects of cartilage regeneration using stem cells. To our knowledge, there have been no published comparisons of the effect of various stem cells for cartilage repair. Therefore, the aim of this study is to investigate which approved cell therapy method is more effective for cartilage repair procedure.

Materials and Methods

Subject

We conducted a retrospective review of the patient to whom underwent cartilage repair surgery using the BMAC and hUCB-MSCs between March 2012 and October 2017. In Korea, the approved indication of hUCB-MSCs (Cartistem®, Medipost Inc.) includes cartilage-defective patient with the International Cartilage Repair Society (ICRS) grade IV lesions, regardless of age or size. BMAC is only approved for patients aged between 15 and 50 years, ICRS grade III to IV, and lesion size 2–10 cm². In this retrospective study, because the indications of the two treatment methods are slightly different, inclusion criteria were set as follows to reduce the selection bias: (1) focal cartilage defects with persistent symptoms of knee joint pain or functional disability, despite a minimum 3 months of conservative treatment; (2) age above 15 years; (3) Kellgren-Lawrence (K-L) grade ≤2; (4) lesion size 2–10 cm²; (5) ICRS grade IV; and (6) minimum 2-year follow-up. In addition, the BMAC group was only for age between 15 and 50 years. Exclusion criteria were as follows: (1) K-L grade ≥3, (2) tricompartment OA, (3) infectious or inflammatory arthropathy, (4) rheumatoid disease of cancer patients, and (5) less than 2-year follow-up. Coexisting knee pathologies such as meniscal tears, meniscus deficiencies, ligament insufficiencies and tibiofemoral axis malalignment were treated during the same surgical procedure.

Finally, total of 52 patients (52 knees, BMAC: 25 knees, BMAC group and hUCB-MSCs: 27 knees, hUCB group) were enrolled in this retrospective study. Detailed demographic data are summarized in Table 1. All patients provided informed consent prior to treatment. This study
protocol was approved by the INHA University Hospital Institutional Review Board (INHA 2019-02-003).

**Surgical Technique**

**BMAC Group.** All the procedures were performed under spinal anesthesia. An arthroscopic examination was performed using a standard anterolateral and anteromedial portal in the lithotomy position. After complete inspection of the joints and assessment of the cartilage defects, marrow aspiration was performed at the ipsilateral iliac crest using an aspiration kit under routine sterile preparation and draping. Sixty milliliters of bone marrow was collected, then centrifuged using a commercially available BMAC system (SmartPReP2®, Harvest Technologies, Plymouth, MA, USA). Coexisting knee pathologies such as meniscus tear (menisectomy or meniscus repair), meniscus deficiency (meniscus allograft transplantation), tibiofemoral axial malalignment (open wedge high tibial osteotomy), and ligamentous insufficiency (anterior cruciate ligament reconstruction) were treated during the same surgery (Table 1). After evaluation and management of coexisting pathologies, a mini-arthrotomy through an incision of approximately 3–4 cm in length was made through the anterolateral or anteromedial portal according to location of the lesion. The chondral defect lesion was debrided and prepared using curettes. Subsequently, small sized holes were made using microfracture awl to the depth about 3–5 mm for the containment of MSCs graft. In the space between the microfracture holes, a 2-mm diameter drill bit was used to create several small holes. After drilling, irrigation was performed to wash out debris. Finally, the lesion area was dried prior to implantation. The defect size was measured and the HA membrane (Hyalofast®, Anika Therapeutics Inc., Bedford, MA, USA) templated according to the size. Subsequently, centrifuged BMAC was added to HA membrane, then implanted on the prepared lesion. After applying the BMAC + HA scaffold, we added fibrin glue (Greenplast®, Greencross, Seoul, Korea) to create more stable fixation of the lesion.

**hUCB Group.** The same procedure was done with BMAC group up to the procedure of managing coexisting pathology and preparing the lesion without marrow aspiration and making containment subchondral hole. Instead of microfracture awl, 5-mm diameter drill bit was used to perform to the depth about 3–5 mm for the containment of MSCs graft. After preparing the lesion, commercially available hUCB-MSCs [Cartistem®, Medipost Inc., composite of hUCB-MSCs 0.5 x 10^7/ml and freeze drying sodium hyaluronate (HA)] were mixed according to the manufacturer’s instructions. After mixing hUCB-MSCs and HA, it becomes a type of gel21. Then, the hUCB-MSCs and HA mixture was implanted into the prepared lesions from the base to the surface slowly to avoid any defects.

After implantation, the knee was then ranged carefully through flexion and extension in order to check the stability of the overlying composite of BMAC or hUCB-MSCs materials. The wound was closed and a long leg splint was applied. All the procedures were performed by the MK Kim. After the procedure, the knee was immobilized for 1 week with an extension knee splint. Continuous passive range of motion exercise was recommended at 1 week after surgery, and weight bearing was restricted for 6 weeks. Partial weight bearing was permitted using a crutch at 6 weeks after surgery, and full weight bearing was allowed at 12 weeks postoperatively.

**Clinical Evaluation**

To assess pain and the functional disability of the knees, we used the visual analogue scale (VAS) pain score (0 = no pain, 10 = worst pain), the International Knee Documentation
Committee (IKDC) subjective score, and the Knee Injury and Osteoarthritis Outcome Score (KOOS)\textsuperscript{25} preoperatively, 6 months, 1 year, and 2 years after surgery.

\textbf{Magnetic Resonance Imaging Evaluation}

Magnetic resonance imaging (MRI) was performed at preoperative, 1 year, and 2 years after surgery. Preoperative and follow-up MRI was performed using a 3.0 T MRI scanner, Discovery MR750W\textsuperscript{®} (GE Healthcare, Chicago, IL, USA) with a dedicated eight channel knee coil. The following sequences were utilized: (a) proton density (PD) spectral presaturation with inversion recovery (SPIR) transversal image [repetition time/echo time (TR/TE), 3,794/28 ms]; field of view (FOV), 160 $\times$ 160 mm; matrix, 480 $\times$ 288; slice thickness (SL), 3.0 mm with 0.3 mm gap; (b) PD SPIR coronal image (TR/TE, 3,513/28.2 ms): FOV, 160 $\times$ 160 mm; matrix, 480 $\times$ 288; SL, 3.0 mm with 0.3 mm gap; (c) T2 SPIR sagittal image (TR/TE, 2,002/71.3 ms): FOV, 160 $\times$ 160 mm; matrix, 288 $\times$ 256; SL, 0.7 mm with 0.3 mm gap; and (d) turbo spin echo T1-weighted sagittal image (TR/TE, 461/15.8 ms): FOV, 160 $\times$ 160; matrix, 480 $\times$ 288; SL, 3.0 mm with 0.33 mm gap.

The MRI images were analyzed using the modified Magnetic Resonance Observation of Cartilage Repair Tissue (M-MOCART) score. Although MRI is unable to accurately determine the status of the cartilage repair\textsuperscript{26}, the M-MOCART (0 = worst cartilage status, 100 = best articular cartilage status) score highly correlated with clinical outcome\textsuperscript{24,27}. The M-MOCART score was classified as excellent if the score was 80 or more and poor outcome when the score was below 50. To avoid bias, a musculoskeletal-trained radiologist who did not participate in the care of patients and was blinded to this study evaluated the MRI images.

\textbf{Second-Look Arthroscopy}

A second-look arthroscopy was conducted 1 year after first surgery in patients who underwent surgical treatment on the same knee for hardware removal and those who agreed to check repaired cartilage condition during a contralateral knee procedure. During the second-look procedures, the repaired cartilage was inspected and evaluated using the ICRS cartilage repair assessment scoring system (score 0–12) including the degree of defect fill, the degree of graft integration to the adjacent normal articular surface, and the gross appearance of the graft surface\textsuperscript{28}.

\textbf{Statistical Analysis}

The repeated-measures analysis of variance test was used to determine the significance of repeated-measures parameters on continuous scale between two groups for metric parameters. The independent Student’s $t$-test was also used to compare the metric parameters between two groups. In a prior power analysis, the number of samples required to perform the noninferiority test between the two groups was calculated using the VAS score, IKDC score, and KOOS score as the reference value ($\alpha = 0.05$, $\beta = 0.2$). The non-inferiority margin of the number of samples was set based on the established minimal clinical important difference (MCID) of VAS score and IKDC subjective score\textsuperscript{29}. Reference values of sample size [mean difference and standard deviation (SD)] and MCID of KOOS score were set based on the previously performed joint cartilage repair research\textsuperscript{10,30}. At least 20 patients in each group were needed. Results of continuous measurements were presented as mean $\pm$ SD, and results on categorical measurements are presented as number. Statistical analysis was conducted using SPSS (Ver 25.0, IBM, Armonk, NY, USA) with significance defined as $P < 0.05$.

\textbf{Results}

\textbf{Clinical Outcomes}

There was a significant difference in age between the two groups due to differences in age indications ($P < 0.0001$). There were no differences in terms of gender, lesion size, number of lesions, or body mass index (BMI) (Table 1). There was a difference in concomitant surgeries between the two groups. The average size of the lesion was 4.33 $\pm$ 1.66 cm$^2$ in the BMAC group and 4.77 $\pm$ 1.81 cm$^2$ in the hUCB group (Table 1).

At final follow-up, VAS scores improved from 5.2 $\pm$ 1.1 to 0.92 $\pm$ 0.98 and from 5.0 $\pm$ 1.2 to 0.85 $\pm$ 0.86 for the BMAC and hUCB groups, respectively (Fig. 1); IKDC subjective scores improved from 44.17 $\pm$ 12.5 to 80.27 $\pm$ 9.48 and 42.02 $\pm$ 13.63 to 81.35 $\pm$ 11.07 for the BMAC and hUCB groups, respectively (Fig. 2). KOOS scores also showed improvement in all categories in both groups.
repair. There was no significant difference between the nearly normal condition (ICRS repair score 8–11) or nearly normal condition (ICRS repair score 8–11) condition.

There were no infections or conversion to total knee arthroplasty (Table 2).

In this study, only 50% of patients (22 patients/44 patients) could be evaluated for the individual findings of previously published studies. The hUCB group showed slightly better results in the M-MOCART score and ICRS repair score; however, these were not statistically significant and there were no significant differences in clinical outcome. To assess cartilage repair status, arthroscopic evaluation is the most accurate method; however, there are limitations for most patients because of the invasive procedure. In this study, only 50% (22 patients/44 patients) could be performed. The use of MRI as an indirect method has been shown to be related to the clinical score.

In this study, we evaluated the degree of cartilage repair status by MRI in patients who did not undergo second-look surgery based on these results. To our knowledge, this was the first study to compare the clinical and MRI outcomes using two types of approved MSCs for cartilage repair.

Two of the patients who underwent BMAC had poor results. One of them was an elite sport coach and one was an active amateur athlete. One patient in the hUCB group, a hard working labor, also showed poor results. Commonly, high physical demand may produce poor results after surgery. In this study, it was difficult to analyze because of the small cases. In a previous study, still under debate, the high physical demand may produce poor results. In this study, it was difficult to analyze because of the small cases. In a previous study, still under debate, the high physical demand may produce poor results.

The demographics of this retrospective study revealed differences between the two groups of subjects, especially in terms of age and concomitant surgery, when treating only patients for whom there were approved indications. Many previous studies have reported that surgical correction of concomitant knee pathology plays a key role in cartilage repair procedure. While biologic and graft failures will occur, the majority of failures were attributed to untreated background factors such as malalignment, meniscal deficiency, and instability. Although the accompanying knee pathologies were different, concomitant surgery has been attempted to adjust for environment favorable for cartilage repair.

**MRI Findings**

At 1-year MRI, there were four patients in the BMAC group and eight patients in the hUCB group with greater than 80 points on M-MOCART score. By contrast, three patients in BMAC and two patients in hUCB had less than 50 points. At 2-year follow-up, there were seven patients in the BMAC group and 10 patients in the hUCB group with better than 80 points on M-MOCART score. By contrast, two patients in the BMAC and one patient in hUCB had less than 50 points. For both year’s M-MOCART scores, there were no significant differences between the groups. Results of M-MOCART score are summarized in Table 2.

**Second-Look Arthroscopy**

Second-look arthroscopy was performed in 12 patients in the BMAC group (Fig. 4) and 16 patients in the hUCB group (Fig. 5) with a mean follow-up of 13.2 months. Eleven patients (91.6%) of BMAC group and 14 patients (87.5%) of hUCB group were assessed as normal (ICRS repair score = 12) or nearly normal (ICRS repair score 8–11) condition repair. There was no significant difference between the groups (P = 0.655) (Table 2).

**Subgroup Analysis Based on Age**

When compared the results of the age-based cartilage repair regardless of the treatment method, there were no significant differences between younger than 45 years (consist of BMAC: 15 cases and hUCB: 2 cases) and older than 45 years (consist of BMAC: 10 cases and hUCB: 25 cases) in MRI outcomes. [1Y-MOCART score (P = 0.849), 2Y-MOCART score (P = 0.83)] and ICRS repair score (P = 0.926). Preoperative and 2-year follow-up, VAS pain score were also no significant difference (Table 3).
cartilage repair. We assumed that the environment of the knee joint in both groups was similarly improved. Generally, cell count and the differential potency of BMAC are reported to decrease with age, and this is the basis for reluctance to use in elderly patients. In fact, there have been reports suggesting that sufficient numbers of MSCs can be collected from elderly patients. Further- more, mononuclear stem cells from younger healthy donors have been reported to differ between donors in terms of differentiation potentials. A recent study have shown that elderly patients underwent successful cartilage repair without significant differences. In the present study, there were no statistical differences between the age groups, regardless of the treatment method, by contrast with the expectation that the cartilage repair effect would be better in younger patients in general.

In the case of hUCB, the same number and capacity of cells could always be transplanted, irrespective of age. However, as mentioned above, BMAC has different cell counts, differentiation potentials, cytokines, and growth factors depending on the age and individual patient’s conditions. Nevertheless, there were no significant differences between the two groups. In this regard, the “medical signaling effect” proposed by Caplan and Correa has attracted attention. Even after implantation of MSCs, the implanted MSCs disappeared after 6–8 weeks, and the cells were replaced with host cells. Therefore, in the cartilage repair procedure, the importance of signaling with the surrounding environment, i.e., paracrine effect, is emphasized more than stem cell count. According to this concept, MSCs, rather than participating in tissue formation, play the role of a site-regulated “drugstore” in vivo by releasing trophic and immunomodulatory factors. Therefore, we believe that even BMAC with a low fraction of stem cell content can achieve effective cartilage repair. Many studies are needed to determine the role of cell count, cytokine, growth factor concentration, and knee joint environment in induction of cartilage repair. Considering the results to date, it would be better to choose the cartilage repair procedure by considering the patient’s individual characteristics. BMAC may be preferred for young, physically healthy people, whereas hUCB-MSCs would be preferred for large sized lesion, old age, or fear for donor site morbidity.

Table 2. The Outcome of MRI (M-MOCART Score), Second-Look Arthroscopy, and Complications of Both Treatment Groups.

|                          | BMAC group (N = 25) | hUCB group (N = 27) | P-value |
|--------------------------|---------------------|---------------------|---------|
| 1Y M-MOCART score       | 65.4 ± 13.46        | 69.63 ± 13.37       | 0.261   |
| 2Y M-MOCART score       | 70.20 ± 13.58       | 73.7 ± 13.2         | 0.351   |
| 1Y 2nd look ICRS repair grade score | 9.42 ± 1.83        | 9.75 ± 2.05         | 0.655   |
| Complication             |                     |                     |         |
| Adhesion                 | 2                   | 3                   |         |
| Infection                | 0                   | 0                   |         |
| TKA conversion           | 0                   | 0                   |         |

BMAC: bone marrow aspirate concentrate; hUCB: human umbilical cord blood; ICRS: The International Cartilage Repair Society; M-MOCART: modified Magnetic Resonance Observation of Cartilage Repair Tissue; MRI: magnetic resonance imaging; TKA: total knee arthroplasty.
The present study has some limitations. First, it is a retrospective study design, with a small sample size, short-term follow-up period, and high risk of bias due attributable to differences in age and coexisting knee pathology between the two treatment groups. Therefore, there is a need for large sample sizes in well-designed prospective randomized trial. Second, biopsy evaluation was not performed. Many studies reported that repaired cartilage was composed of hyaline like cartilage\textsuperscript{30,44,45}. However, as Gobbi et al.\textsuperscript{31} found, although the ICRS repair score was high, it may not be possible to show precise results because there were cases where the actual biopsy showed a mixed (hyaline + fibrocartilage) repair. In this case, we could not guarantee the long-term clinical results. Furthermore, the second-look arthroscopy was performed at 1 year after surgery; therefore, we could not determine whether the repaired cartilage changed after the first year. Also, second-look arthroscopy was conducted only with a small patient in both groups. There is a high risk of selection bias to evaluate outcomes. Third, there are a variety of factors that can affect the results of cartilage repair, including age, BMI, coexisting knee lesions, and physical demand, all of which may limit the interpretation of the results. Fourth, the use of fibrin glue in the BMAC group may have affected the outcomes. According to a literature review, the fibrinogen concentration can affect cell proliferation and migration\textsuperscript{46,47}.

Despite these limitations, our present study has some merits. To our knowledge, this was the first study to compare the clinical and MRI outcomes using BMAC and hUCB-MSCs for cartilage repair. This can help surgeons determine what kind of cartilage repair method is better for their patients. We also compared the results of the age-based cartilage repair regardless of the treatment method. In this way, we were able to suggest what additional research would be needed to determine effective cartilage repair in the elderly while supporting existing research results.

Figure 4. A representative case of cartilage repair using BMAC (female/47-year-old). (A) Standing anteroposterior knee radiography showed varus alignment of left knee (HKA axis: varus 7°). (B and C) Preoperative MRI exam showed ICRS grade IV lesion (red arrow) through T2 proton density fat suppression (T2 PD FS) at MFC through sagittal (B), coronal view (C). (D, E, F) Intraoperative arthroscopic findings. A 4.5 cm² sized ICRS IV lesion was observed at MFC through arthroscope (D). After preparation for implantation (E), BMAC + HA scaffold implantation was performed with additional fibrin glue fixation (F). (G, H) After 1-year follow-up, standing anteroposterior radiography showed corrected varus alignment (HKA axis: valgus 0.8°) (G). During second-look arthroscopy evaluation, repaired cartilage had some overgrowth, but were well integrated to surrounding tissues and smoothed surface (H). (I, J) At 2-year follow-up, repaired cartilage (yellow arrow) completely filled defect, showed smooth surface and well integrated to around tissue at T2 PD FS MRI images. There was no definite subchondral edema at MRI exam sagittal (I), and coronal (J) view. BMAC: bone marrow aspirate concentrate; HKA: Hip-Knee-Ankle; ICRS: The International Cartilage Repair Society; MFC: medial femoral condyle; and MRI: magnetic resonance imaging.
Conclusion

The results of our study are encouraging and show that implantation of MSCs from BMAC or hUCB-MSCs for cartilage repair in knee osteoarthritis is safe and effective for repairing cartilage lesion in each indication. Therefore, if applied with proper patient indication, both methods could be expected satisfactory outcome. However, due to differences in patient groups, the effectiveness of the two methods could not be accurately compared. Thus, large cases and a well-controlled prospective design with long-term follow-up studies are needed to set proper indication.

Ethical Approval

All procedures performed in this study involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards. The protocol of this study was approved by the Inha University Hospital Institutional Review Board (approval number: INHAUH 2019-02-003).

Statement of Human and Animal Rights

All procedures in this study were conducted in accordance with the Korea Food and Drug Administration (KFDA) guideline for cartilage repairing procedure. And the protocol of this study was...
approved by the Inha University Hospital Institutional Review Board (approval number: INHAUH 2019-02-003).

**Statement of Informed Consent**

Informed consent was obtained from all individual participants included in the study.

**Declaration of Conflicting Interests**

The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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**References**

1. Alford JW, Cole BJ. Cartilage restoration, part 1: basic science, historical perspective, patient evaluation, and treatment options. Am J Sports Med. 2005;33(2):295–306.
2. Scotti C, Gobbi A, Karnatzikos G, Martin I, Shimomura K, Lane JG, Peretti GM, Nakamura N. Cartilage repair in the inflamed joint: considerations for biological augmentation toward tissue regeneration. Tissue Eng Part B Rev. 2016;22(2):149–159.
3. Gobbi A, Karnatzikos G, Kumar A. Long-term results after microfracture treatment for full-thickness knee chondral lesions in athletes. Knee Surg Sports Traumatol Arthrosc. 2014;22(9):1986–1996.
4. Beris AE, Lykissas MG, Kostas-Agnantis I, Manoudis GN. Treatment of full-thickness chondral defects of the knee with autologous chondrocyte implantation: a functional evaluation with long-term follow-up. Am J Sports Med. 2012;40(3):562–567.
5. Orth P, Rey-Rico A, Venkatesan JK, Madry H, Cucchiari M. Current perspectives in stem cell research for knee cartilage repair. Stem Cells Cloning. 2014;7:1–17.
6. Park Y-B, Ha C-W, Rhim JH, Lee H-J. Stem cell therapy for articular cartilage repair: review of the entity of cell populations used and the result of the clinical application of each entity. Am J Sports Med. 2018;46(10):2540–2552.
7. Robinson PG, Murray IR, West CC, Goudie EB, Yong LY, White TO, LaPrade RF. Reporting of mesenchymal stem cell preparation protocols and composition: a systematic review of the clinical orthopaedic literature. Am J Sports Med. 2019;47(4):991–1000.
8. Piuzzi NS, Hussain ZB, Chahla J, Cinque ME, Moatshe G, Mantripragada VP, Muschler GF, LaPrade RF. Variability in the preparation, reporting, and use of bone marrow aspirate concentrate in musculoskeletal disorders: a systematic review of the clinical orthopaedic literature. J Bone Joint Surg Am. 2018;100(6):517–525.
9. Chahla J, Dean CS, Moatshe G, Pascual-Garrido C, Serra Cruz R, LaPrade RF. Concentrated bone marrow aspirate for the treatment of chondral injuries and osteoarthritis of the knee: a systematic review of outcomes. Orthop J Sports Med. 2016;4(1):2325967115625481.
10. Gobbi A, Whyte GP. One-stage cartilage repair using a hyaluronic acid-based scaffold with activated bone marrow-derived mesenchymal stem cells compared with microfracture: five-year follow-up. Am J Sports Med. 2016;44(11):2846–2854.
11. Samsonraj RM, Rai B, Sathiyanathan P, Puan KJ, Rötzschke O, Hui JH, Raghunath M, Stanton LW, Nurcombe V, Cool SM. Establishing criteria for human mesenchymal stem cell potency. Stem Cells. 2015;33(6):1878–1891.
12. Baxter MA, Wynn RF, Jowitt SN, Fairbairn LJ, Bellantuono I. Study of telomere length reveals rapid aging of human marrow stromal cells following in vitro expansion. Stem Cells. 2004;22(5):675–682.
13. Fortier LA, Potter HG, Rickey EJ, Schnabel LV, Foo LF, Chong LR, Stokol T, Cheetham J, Nixon AJ. Concentrated bone marrow aspirate improves full-thickness cartilage repair compared with microfracture in the equine model. J Bone Joint Surg Am. 2010;92(10):1927–1937.
14. Krych AJ, Nawabi DH, Farshad-Amacker NA, Jones KJ, Maak TG, Potter HG, Williams RJ. Bone marrow concentrate improves early cartilage phase maturation of a scaffold plug in the knee: a comparative magnetic resonance imaging analysis to platelet-rich plasma and control. Am J Sports Med. 2016;44(1):91–98.
15. Cotter EJ, Wang KC, Yanke AB, Chubinskaya S. Bone marrow aspirate concentrate for cartilage defects of the knee: from bench to bedside evidence. Cartilage. 2018;9(2):161–170.
16. Erices A, Conget P, Minguell JJ. Mesenchymal progenitor cells in human umbilical cord blood. Br J Haematol. 2000;109(1):235–242.
17. Berg L, Koch T, Heerkens T, Bessonov K, Thomsen P, Betts D. Chondrogenic potential of mesenchymal stromal cells derived from equine bone marrow and umbilical cord blood. Vet Comp Orthop Traumatol. 2009;22(5):363–370.
18. Futami I, Ishijima M, Kaneko H, Tsuji K, Ichikawa-Tomikawa N, Sadatsuki R, Muneta T, Arikawa-Hirasawa E, Sekiya I, Kaneko K. Isolation and characterization of multipotential mesenchymal cells from the mouse synovium. PLoS One. 2012;7(9):e45517.
19. Sakaguchi Y, Sekiya J, Yagishita K, Muneta T. Comparison of human stem cells derived from various mesenchymal tissues: superiority of synovium as a cell source. Arthritis Rheum. 2005;52(8):2521–2529.
20. Weiss ML, Anderson C, Medicetty S, Sheshareddy KB, Weiss RJ, VanderWerff I, Troyer D, McIntosh KR. Immune properties of human umbilical cord Wharton’s jelly-derived cells. Stem Cells. 2008;26(11):2865–2874.
21. Park Y-B, Ha C-W, Lee C-H, Yoon YC, Park Y-G. Cartilage regeneration in osteoarthritic patients by a composite of...
allogeneic umbilical cord blood-derived mesenchymal stem cells and hyaluronate hydrogel: results from a clinical trial for safety and proof-of-concept with 7 years of extended follow-up. Stem Cells Transl Med. 2017;6(2):613–621.

22. Koh Y-G, Kwon O-R, Kim Y-S, Choi Y-J, Tak D-H. Adipose-derived mesenchymal stem cells with microfracture versus microfracture alone: 2-year follow-up of a prospective randomized trial. Arthroscopy. 2016;32(1):97–109.

23. Koh Y-G, Choi Y-J, Kwon S-K, Kim Y-S, Yeo J-E. Clinical results and second-look arthroscopic findings after treatment with adipose-derived stem cells for knee osteoarthritis. Knee Surg Sports Traumatol Arthrosc. 2015;23(5):1308–1316.

24. Kim YS, Choi YJ, Lee SW, Kwon OR, Suh DS, Heo DB, Koh YG. Assessment of clinical and MRI outcomes after mesenchymal stem cell implantation in patients with knee osteoarthritis: a prospective study. Osteoarthr Cartil. 2016;24(2):237–245.

25. Hambly K, Griva K. IKDC or KOOS? Which measures symptoms and disabilities most important to postoperative articular cartilage repair patients? Am J Sports Med. 2008;36(9):1695–1704.

26. de Windt TS, Welsch GH, Brittberg M, Vonk LA, Marlovits S, Trattning S, Saris DBF. Is magnetic resonance imaging reliable in predicting clinical outcome after articular cartilage repair of the knee? A systematic review and meta-analysis. Am J Sports Med. 2013;41(7):1695–1702.

27. Welsch GH, Mamisch TC, Zak L, Blanke M, Olk A, Marlovits S, Trattning S. Evaluation of cartilage repair tissue after matrix-associated autologous chondrocyte transplantation using a hyaluronic-based or a collagen-based scaffold with morphological MOCART scoring and biochemical T2 mapping: preliminary results. Am J Sports Med. 2010;38(5):934–942.

28. Brittberg M, Winalski CS. Evaluation of cartilage injuries and repair. J Bone Joint Surg Am. 2003;85(A Suppl 2):58–69.

29. Collins NJ, Misra D, Felson DT, Crossley KM, Roos EM. Measures of knee function. Arthritis Care Res (Hoboken). 2011;63(0 11): S208–S228.

30. Gobbi A, Karnatzikos G, Sankineani SR. One-step surgery with multipotent stem cells for the treatment of large full-thickness chondral defects of the knee. Am J Sports Med. 2014;42(3):648–657.

31. Gobbi A, Scotti C, Karnatzikos G, Mudigere A, Castro M, Peretti GM. One-step surgery with multipotent stem cells and Hyaluronan-based scaffold for the treatment of full-thickness chondral defects of the knee in patients older than 45 years. Knee Surg Sports Traumatol Arthrosc. 2017;25(8):2494–2501.

32. Guermazi A, Roemer FW, Alizai H, Winalski CS, Welsch G, Brittberg M, Trattning S. State of the art: MR imaging after knee cartilage repair surgery. Radiology. 2015;277(1):23–43.

33. Blackman AJ, Smith MV, Flanagan DC, Matava MJ, Wright RW, Brophy RH. Correlation between magnetic resonance imaging and clinical outcomes after cartilage repair surgery in the knee: a systematic review and meta-analysis. Am J Sports Med. 2013;41(6):1426–1434.

34. Krych AJ, Hevesi M, Desai VS, Camp CL, Stuart MJ, Saris DBF. Learning from failure in cartilage repair surgery: an analysis of the mode of failure of primary procedures in consecutive cases at a tertiary referral center. Orthop J Sports Med. 2018;6(5):2325967118773041.

35. Sharma L, Song J, Felson DT, Cahue S, Shamiyeh E, Dunlop DD. The role of knee alignment in disease progression and functional decline in knee osteoarthritis. JAMA. 2001;286(2):188–195.

36. Payne KA, Didiano DM, Chu CR. Donor sex and age influence the chondrogenic potential of human femoral bone marrow stem cells. Osteoarthr Cartil. 2010;18(5):705–713.

37. Bodle JC, Teeter SD, Hluck BH, Hardin JW, Bernacki SH, Loboa EG. Age-related effects on the potency of human adipose-derived stem cells: creation and evaluation of superlot and implications for musculoskeletal tissue engineering applications. Tissue Eng Part C Methods. 2014;20(12):972–983.

38. Juneca S, Viswanathan S, Ganguy M, Veillette C. A simplified method for the aspiration of bone marrow from patients undergoing hip and knee joint replacement for isolating mesenchymal stem cells and In Vitro chondrogenesis. Bone Marrow Res. 2016;2016:3152065.

39. Caplan AI, Correa D. The MSC: an injury drugstore. Cell Stem Cell. 2011;9(1):11–15.

40. Park WS, Ahn SY, Sung SI, Ahn J-Y, Chang YS. Strategies to enhance paracrine potency of transplanted mesenchymal stem cells in intractable neonatal disorders. Pediatr Res. 2018;83(1–2):214–222.

41. van Poll D, Parekkadan B, Cho CH, Berthiaume F, Nahmias Y, Tilles AW, Yarmush ML. Mesenchymal stem cell-derived molecules directly modulate hepatocellular death and regeneration in vitro and in vivo. Hepatology. 2008;47(5):1634–1643.

42. Prockop DJ, Kota DJ, Bazhanov N, Reger RL. Evolving paradigms for repair of tissues by adult stem/progenitor cells (MSCs). J Cell Mol Med. 2010;14(9):2190–2199.

43. Yanke AB, Chubinskaya S. The state of cartilage regeneration: current and future technologies. Curr Rev Musculoskelet Med. 2015;8(1):1–8.

44. Gigante A, Ceconi S, Calcagno S, Busilacchi A, Enea D. Arthroscopic knee cartilage repair with covered microfracture and bone marrow concentrate. Arthrosc Tech. 2012;1(2):e175–e180.

45. Enea D, Ceconi S, Calcagno S, Busilacchi A, Manzotti S, Kaps C, Gigante A. Single-stage cartilage repair in the knee with microfracture covered with a resorbable polymer-based matrix and autologous bone marrow concentrate. Knee. 2013;20(6):562–569.

46. Guo H-D, Wang H-J, Tan Y-Z, Wu J-H. Transplantation of marrow-derived cardiac stem cells carried in fibrin improves cardiac function after myocardial infarction. Tissue Eng Part A. 2011;17(1–2):45–58.

47. Lee HH, Haleem AM, Yao V, Li J, Xiao X, Chu CR. Release of bioactive adeno-associated virus from fibrin scaffolds: effects of fibrin glue concentrations. Tissue Eng Part A. 2011;17(15–16):1969–1978.