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Does progress in microfracture techniques necessarily translate into clinical effectiveness?

Muthu S *et al.* Effectiveness of Microfracture techniques

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Abstract

BACKGROUND

Multitudinous advancements have been made to the traditional microfracture (MFx) technique, which have involved delivery of various acellular 2nd generation MFx and cellular MFx-III components to the area of cartilage defect. The relative benefits and pitfalls of these diverse modifications of MFx technique are still not widely understood.

AIM

To comparatively analyze the functional, radiological, and histological outcomes, and complications of various generations of MFx available for the treatment of cartilage defects.

METHODS

A systematic review was performed using PubMed, Embase, Web of Science, Cochrane, and Scopus. Patients of any age and sex with cartilage defects undergoing any form of MFx were considered for analysis. We included only randomized controlled trials (RCTs) reporting functional, radiological, histological outcomes or complications of various generations of MFx for the management of cartilage defects. Network meta-analysis (NMA) was conducted in Stata and Cochrane's Confidence in NMA (CINeMA) approach was utilized for appraisal of evidence.

RESULTS

A systematic review was performed using PubMed, Embase, Web of Science, Cochrane, and Scopus. Patients of any age and sex with cartilage defects undergoing any form of MFx were considered for analysis. We included only RCTs reporting functional, radiological, histological outcomes or complications of various generations of MFx for the management of cartilage defects. NMA was conducted in Stata and Cochrane's CINeMA approach was utilized for appraisal of evidence.

CONCLUSION

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Key Words: Cartilage injury; Microfracture; Mesenchymal stem cells; Platelet-rich plasma; Bone marrow aspiration concentrates; Clinical outcome; Radiological outcome; Meta-analysis; Network meta-analysis

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Core Tip: Chondral lesions have been reported in 60% of patients undergoing arthroscopic procedures of the knee; and such defects are described as one of the leading causes of chronic knee pain. As compared with the other cartilage restoration strategies, microfracture (MFx) is relatively cost-effective, simple, minimally-invasive and may also be performed in a single stage. Nevertheless, recent studies have demonstrated that modifications of the traditional MFx technique, such as the use of

synthetic and autologous biological adjuvants may enhance the repair tissue quality, resilience, and overall efficacy of the procedure. Based on the current network meta-analysis we could conclude that the use of acellular and cellular adjuvants has shown only marginal improvement in the clinical (pain and functional scores) and radiological outcome in patients undergoing microfracture for cartilage defects of the knee. The safety and efficacy of the higher generation MFx procedures are also clearly evident from our review. However, there is a substantial potential for further improvement in the cellular components (chondrocytes over other cellular lineage), culture or processing methodology, delivery modalities (including appropriate scaffolds); as well as better surgical techniques to achieve demonstrable significant outcome improvement.

INTRODUCTION

Lesions of the articular cartilage of the knee remain a challenging clinical entity in view of the limited capacity of the cartilaginous tissues to heal and potential progression to chronic degenerative arthritis^[1]. The deficient endogenous cartilage repair mechanism has been attributed to the poor recruitment of regenerative cells into the area of cartilage defect^[2]. Based upon the theory of marrow stimulation by subchondral drilling purported by Pridie *et al* (1959), Steadman *et al*^[3] popularized the concept of microfracture (MFx) technique, whereby the migration of the growth factors and mesenchymal stem cells (MSCs) across the subchondral bone stimulates the development of the hyaline-like fibrocartilage. As compared with the other cartilage restoration strategies, MFx is relatively cost-effective, simple, minimally-invasive and may also be performed in a single stage^[4]. Despite still being regarded as the gold-standard first-line treatment for cartilage deficiencies of the knee, there are concerns regarding their long-term outcomes and durability of the restored fibrocartilage^[5,6]. In this context, alternate cartilage restoration procedures such as autologous chondrocyte implantation (ACI), osteoarticular transfer system and osteochondral allograft transplantation have been advocated as the better treatment strategies in the recent

years. In fact, the United Kingdom² National Institute for Health and Care Excellence, in a recent assessment, has recommended for the abandonment of MFx in favor of ACI in the management of articular knee defects^[7-11].

Nevertheless, recent studies have demonstrated that modifications of the traditional MFx technique, such as² the use of synthetic and autologous biological adjuvants may enhance the repair tissue quality, resilience and overall efficacy of the procedure^[7,11]. Some researchers have purported that the suboptimal efficacy of the traditional marrow stimulating techniques may be attributed to the insufficient concentrations of² MSCs and growth factors getting released from subchondral marrow. To circumvent this limitation, it has been proposed that supplementation of MFx with intra-articular adjuvants in the form of platelet-rich plasma (PRP) or hyaluronic acid (HA) can improve the outcome^[12-18]. In addition, augmentation of defect with scaffolding matrix or cell-free polymer-based implant can provide a bioreactor-like structure, over which the marrow elements get trapped, concentrated and thereby, facilitate the restoration of an effective cartilage layer^[19-21]. MFx has also been combined with diverse cellular additives like bone marrow aspiration concentrates (BMAC), MSCs, and peripheral blood stem cells (PBSCs). While individual studies on these biological augmentation [popularly described as “microfracture plus” (MFx+)] techniques have demonstrated encouraging histological and clinical outcomes, our understanding regarding these² techniques has been limited by substantial heterogeneity among the study cohorts and paucity of high quality, prospective trials.

The purpose of our study was to consolidate the available evidence; compare the clinical, functional and radiological outcomes of three different generations of MFx techniques (traditional MFx, MFx + acellular additives, and MFx + cellular additives); and to provide the best recommendations on their relative efficacies, advantages, complications and pitfalls in the management of cartilaginous defects of the knee joint.

MATERIALS AND METHODS

Extraction of data

Cochrane Consumers and Communication Group recommendations were followed for data extraction from the included studies. The following were extracted, and a master chart was prepared: (1) Study characteristics: Author name, country, publication year, number of patients in the study; (2) baseline characteristics: Age for the individual treatment arms, gender proportions, cartilage defect size, interventions analyzed, and duration of follow-up; (3) functional outcomes: Visual Analog Scale (VAS) score for pain, Western Ontario McMaster Universities Osteoarthritis Index score, Tegner score, Lysholm score, International Knee Documentation Committee (IKDC) score, Cincinnati score, and Knee Osteoarthritis Outcome Scale (KOOS) score; (4) radiological outcomes: Magnetic resonance observation of cartilage repair tissue (MOCART) score, and successful magnetic resonance imaging (MRI)-based defect filling ($\geq 2/3^{\text{rd}}$ of the defect); and (5) complications: Adverse events and failures (patient requiring revision surgeries).

Data extraction was performed independently by two reviewers. The different generations of MFx techniques, described in accordance with the ORG classification, include: First-generation MFx (MFx-I) representing the traditional MFx technique; second-generation MFx (MFx-II) involving MFx-I combined with acellular additives [such as PRP, HA, collagen, and procedures such as autologous matrix-induced chondrogenesis (AMIC)]; and third-generation MFx (MFx-III) involves combining MFx-I with cellular additives such as MSCs, BMAC, PBSCs, and stromal vascular fraction (SVF)^[24].

We anticipated heterogeneity among the diverse studies in the duration of follow-up for the analysis of outcome measures. Therefore, we analyzed individual outcomes at short-term (1 years and 2 years), intermediate-term (5 years), and if available long-term (≥ 10 years), based on the available data at individual time points for the outcome concerned. The risk of bias of included studies was analyzed RoB2 tool from Cochrane group^[25]. It was agreed upon that studies with a high risk of bias would be excluded from the study.

RESULTS

Overall, 9416 articles were shortlisted for initial screening. De-duplication resulted in 3584 articles. Title and abstract screening excluded 3231 articles. Among them, 353 articles qualified for full-text review; and 44 eligible RCTs^[4,9,13,15,19,20,31-68] with 2629 included patients qualified for inclusion in the study. PRISMA flow diagram for the inclusion of studies is shown in Figure 1.

The included studies reported at least one of the generations of MFx employed in cartilage defect management. The baseline characteristics of the studies included in the network are presented in Table 1. Norway ($n = 6$), Germany ($n = 5$), and United States ($n = 5$) were the leading countries reporting the highest number of RCTs in the field. The network plot has been presented in Figure 1. The network had 36 possible pair-wise comparisons, among which, 14 had direct evidence data. The network had 42 two-armed studies and 2 multi-armed studies. We did not find significant variability among the characteristics of the included patients in the network concerning age and gender proportions. The mean age of the patients included in the trials was 39.40 (± 9.46) years. The mean follow-up in the included trials ranged between 1 and 15 years.

Quality assessment

None of the included studies demonstrated high risk of bias to warrant exclusion from the study. The risk of bias in the pairwise comparisons is presented in Supplementary Figure 1. We did not find any significant publication bias using the funnel plot for most of the outcome measures analyzed. When publication bias was noted, we adjusted using the “trim and fill” method to identify the missing studies and their effects on the overall estimate. We did not find any significant impact of the missing studies on the overall outcomes, as shown in Supplementary Figure 2.

Network analysis results

We performed a pooled NMA using a frequentist approach to every outcome of interest. Among all the treatment arms in the network, MFx-I had high data strength as

compared with all the other comparators (as shown in the network plots in Supplementary Figure 3). Therefore, MFX-I is taken as the constant comparator and all the outcomes have been reported in comparison to the performance of MFX-I. The outcomes have been analyzed in terms of pain, functional outcomes, radiological outcomes, adverse effects, and failures.

Pain: Inference from the VAS score is taken into consideration for pain outcomes. VAS score was reported at one year in 13 studies^[4,15,33,38,41,44,45,49,53,55-58] involving 676 patients, at two years in 10 studies^[4,15,33,38,41,45,50,53,57,68] involving 690 patients and at 5 years in 3 studies^[39,41,54] involving 297 patients. The pooled forest plot of the VAS score outcome based on the aforementioned follow-up time points is presented in Figures 2, 4, and 5 respectively. Although we did not note a statistically significant improvement in the pain reduction with the advancements to the traditional MFX, the SUCRA ranking of the interventions were consistent in favouring the higher generations in the following order MFX-III > MFX-II > MFX-I as shown in Table 2.

Functional outcomes: The functional outcomes were reported using KOOS, Lysholm score, IKDC score, and Cincinnati score. Figure 2 shows the pooled forest plot of various scores. KOOS score was reported at one year in 8 studies^[32,33,44,46,51,55-57] involving 569 patients, and at 2 years in 4 studies^[32,33,51,57] involving 361 patients. Lysholm score was reported at 1 year in 10 studies^[4,33,35,41,44,47,48,53,59,65] involving 499 patients, and at 2 years in 8 studies^[4,15,33,39,41,47,53,59] involving 516 patients. IKDC score was reported at 1 year in 15 studies^[15,35,37,43-45,56-60,64,66,67] involving 631 patients, at 2 years in 13 studies^[15,37,39,43,45,50,57-59,64,66-68] involving 782 patients, and at 5 years in 4 studies^[39,54,58,59] involving 295 patients. Cincinnati score was reported at 1 year in 3 studies^[31,38,65] involving 117 patients, and at 2 years in 4 studies^[31,38,39,50] involving 349 patients.

The functional outcomes reported at 1, 2, and 5-year time points using the aforementioned scores were clubbed together for the sake of understanding (despite the

limitation of such an approach), in view of the heterogeneity in the reporting of functional outcomes among the reviewed studies.

One-year functional outcomes: The pooled forest plot of the functional outcomes, subgrouped based on the individual scores at 1 year, is presented in Figure 2. We observed statistically significant outcome in the higher generations of MFX evaluated with IDKC score (WMD = 3.40; 95%CI: 0.65, 6.16; $P = 0.045$; without significant heterogeneity). However, the difference was not clinically relevant; and less than the minimum clinical difference for the outcome concerned. Although we did not note a statistically significant improvement in most of the functional outcomes with the advancements to the traditional MFX; we observed that (with the exception of Lysholm score) the SUCRA ranking of the interventions consistently favoured the higher generations in the following order: MFX-III > MFX-II > MFX-I (Table 2).

Two-year functional outcome: The pooled forest plot of the functional outcomes, subgrouped based on the individual scores at 2 years, is presented in Figure 4. We did not note statistically significant difference with the higher generations of MFX with regard to the functional scores such as KOOS, Lysholm score, IDKC score, and Cincinnati score. Nevertheless, similar to the functional outcome at 1-year time point; SUCRA rankings of interventions were consistent in favouring the higher generations in the following order MFX-III > MFX-II > MFX-I (for all outcome measures except the Lysholm score (Table 2).

Five-year functional outcomes: We did not have sufficient data points to evaluate mid-term and long-term functional outcomes. However, based on the available data, there was no significant change in the functional outcome with the higher generations of MFX, as compared to the traditional technique (based on IKDC score; Figure 5). Nevertheless, as with the earlier time points, the SUCRA ranking of interventions favoured the higher generations (in the order MFX-III > MFX-I; Table 2).

Radiological outcomes

The MOCART (magnetic resonance observation of cartilage repair tissue) Score and MRI defect filling ($> 2/3^{\text{rd}}$) have been used to report the radiological outcomes in the included studies. The MOCART score was reported at 1 year in 8 studies^[4,32,44,56,57,59,60,65] involving 439 patients, and at 2 years in 3 studies^[13,32,59] involving 230 patients. The MRI-based defect filling was reported at 1 year in 17 studies^[19,20,31,37,38,40,43-45,47,56,57,60,62-64,67] involving 847 patients, and at 2 years in 10 studies^[13,19,31,38,45,47,50,64,67,68] involving 610 patients.

The pooled forest plots of the radiological outcomes, sub-grouped based on the individual scores at 1- and 2-year time points, are presented in Figures 3 and 4, respectively. We observed statistically better MOCART score in the higher generations of MFx (WMD = 17.44; 95%CI: 0.72, 34.16; $P = 0.025$; without significant heterogeneity) at 1 year. However, the difference was not maintained at 2 years. Although we did not note a statistically significant improvement in the MRI-filling with the advancements to the traditional MFx, the SUCRA ranking of the interventions were consistent in favouring the higher generations in the following order MFx-III $>$ MFx-II $>$ MFx-I (Table 2).

Complications

Adverse events: The adverse events following the compared interventions were reported in 32 studies^[9,19,20,31-33,37-39,43,44,46-48,50-55,57,58,60-63,65-67,69-75] involving 1752 patients. Figure 3 shows the pooled forest plot of the reported complications for the analyzed interventions. In comparison with MFx-I, there was no statistically significant difference in the reported rates of adverse events in the higher generations. On the other hand, the SUCRA ranking of the interventions favoured the higher generations in the following order MFx-III $>$ MFx-II $>$ MFx-I (Table 2); thereby, highlighting the safety of the higher generations in comparison with the traditional technique.

Failures: The need for subsequent procedures following the interventions was considered as treatment failure, and the same was reported in 31 studies^[4,31,33,34,38-42,46,48,57,59,61,63-65,69,72,73,76,77] involving 1059 patients. Figure 3 shows the pooled forest plot of the failure events for the reported interventions. In comparison with MFX-I, there was no statistically significant difference in the failure events among the higher generations of MFX techniques. Moreover, the SUCRA ranking of the interventions favoured the higher generations in the following order MFX-III > MFX-II > MFX-I (Table 2); thus, highlighting the reliability of the higher generations in comparison to the traditional technique.

Sensitivity & ¹subgroup analysis

We did not observe significant heterogeneity across various outcomes analyzed in the network (based upon the heterogeneity values in the corresponding individual forest plots of pairwise comparisons of interventions). We sub-grouped and analyzed the studies based on the outcome measures and follow-up time point in order to avoid any heterogeneity in the pooled results.

¹Consistency

We did not observe any significant evidence of global inconsistency, which could have affected the transitivity of the network results. The consistency analysis was performed for the individual outcomes; and the ¹chi-square values in the corresponding pair-wise comparison forest plots were presented. We noted the indirect pooled estimates to have wider CI compared to direct estimates in some of the paired networks analysed (although ¹without any evidence of systematic differences concerning the potential effect modifiers). We considered these apparent inconsistencies to be the effect of true differences between the direct and indirect estimates. The indirect estimates were considered to reflect a more precise estimate, since they were from a network involving a larger number of studies.

Confidence in evidence

Upon grading the paired comparisons in the network using the CINeMA approach, a “high” confidence was noted across a majority of the paired comparisons (Table 3). However, some of the comparison pairs demonstrated “moderate” confidence. The lack of precision was the most common reason, which downgraded the quality of evidence in the indirect estimates, in view of wider CIs extending on either side of the axes. We also observed some concerns due to certain “within-study bias”, following selective reporting of some of the outcome measures.

DISCUSSION

Chondral lesions have been reported in 60% of patients undergoing arthroscopic procedures of the knee; and such defects are described as one of the leading causes of chronic pain^[78-81]. These defects may result from acute trauma, repetitive microtrauma, osteochondritis dissecans or early osteoarthritis; and can produce symptoms like pain, swelling, catching, stiffness and locking^[33]. In 1743, Hunter *et al*^[82,83] described the challenge of cartilaginous injury by stating that, “once the cartilage is destroyed, it never recovers”. These observations still hold true; and the avascular as well as aneural nature of cartilage substantially limits its ability to self-regenerate^[84]. If left untreated, a transgressed cartilage gradually results in severe osteoarthritis of the joint and ensuing long-standing disability^[85].

Superficial cartilage deficiencies do not induce a local inflammatory response; therefore, despite proliferation of matrix molecules and chondrocytes, the surface is not adequately restored^[86]. When the cartilage defect penetrates the subchondral plate, the vascularized bone marrow can enable the formation of clot rich in chondroprogenitor cells, fibrin and bioactive molecules; which in turn, facilitates the formation of type I collagen and fibrocartilage^[87]. This is the rationale underlying the MFX technique, which has traditionally remained the first-line treatment for small to medium-sized defects^[88]. The purported benefits of the procedure include low cost, easy technique and proven improvement in short-term outcome^[87,88]. Nevertheless, 47% to 80% of patients have

been reported to demonstrate substantial functional deterioration at 18 to 36 months post-surgically^[10], which may be attributed to the poor viscoelastic properties of the restored fibrocartilage^[89]. Since the initial description of MFX technique, multitudinous attempts have been made in the fields of tissue engineering and cartilage repair in an attempt to find the “holy grail”, which enables the restoration of hyaline cartilage that can consistently integrate into the deficiency^[42].

Evolution of MFX

In the traditional MFX technique described by Steadman *et al*^[3], the debridement of the unstable cartilaginous tissues is initially performed arthroscopically; and a well-shouldered vertical wall is created around the periphery of the lesion. Following this, layers of calcified cartilage are removed using a curette. An arthroscopic awl is then utilized in a direction perpendicular to the bone in order to create holes in the subchondral plate around 3-4 mm apart (ascertaining that the interposed subchondral bone between the MFX perforations is maintained intact). Alternately, microdrilling using a 1.5 mm drill may be performed to perforate the subchondral plate to a depth of 1 cm.

While lesions smaller than 2 cm² in low-demand individuals are amenable to treatment with traditional MFX technique; lesions larger than 4cm² have been purported to require additional adjuvant modalities too^[90]. Diverse acellular biomaterials such as alginate, collagen, tri-copolymer and poly-lactic-glycolic acid have been utilized for engineering of cartilaginous tissues^[91]. These tissues serve as carriers for delivery of cells and growth factors; as well as provide an appropriate milieu for tissue regeneration^[92].

The cell therapy for cartilage repair was initially proposed by Robert Langer and Charles Vacanti in the 1980s using the technology of tissue engineering^[93]; and cellular therapeutic innovation was eventually realized in 1994, when Brittberg *et al*^[94] described the ACI technique. Further on, scaffold-based ACI (matrix-induced ACI-MACI: FDA-approved in 2016) technique has also been described as a modification of the traditional

3 MFx. The discovery of adult stem cells resulted in a paradigm shift in the field of regenerative medicine^[95]. A variety of stem cell-based therapies involving multipotent MSCs implantation (like bone marrow, adipose tissue, synovium, periosteum, peripheral blood, *etc.*) have been employed for cartilage repair. The chondrogenesis and development of neo-cartilaginous tissues from such undifferentiated 3 MSCs can be guided using growth factors, and other biophysical or biomechanical stimuli^[96,97].

As an alternative form of cell-based therapy, Gobbi *et al*^[10] described the technique of implanting the bone marrow aspirate concentrate delivered *via* HA-based scaffold (HA-BMAC) over the micro-fractured area. Such an approach relies on the presence of MSCs and growth factors at the deficient zone so as to steer chondrogenesis. They concluded that such an approach yielded successful medium-term clinical outcome with restoration of durable cartilage, irrespective of the size and age of the lesion.

Despite such extensive publications, there has been a substantial dearth of large-scale, high-quality RCTs on this subject. In a recent systematic review; among 540 reviewed manuscripts, only 10 studies were found to be methodologically sufficient to be included for final analysis. The current evidence on this subject is therefore, still largely unclear^[98]. The purpose of the current NMA was to comprehensively analyse the existing literature on chondral injuries of the knee; and comparatively evaluate the histological, radiological and clinical outcome following 3 different generations of MFx, namely traditional MFx (MFx-I), modified MFx technique using acellular adjuvant (MFx-II); and modified MFx technique using cellular adjuvant (MFx-III).

Observations from our study

Clinical and functional outcome: Overall, in our meta-analysis, we compared the pain scores and functional outcome measures (KOOS, Lysholm score, IKDC score, and Cincinnati scores) among the three generations of MFx. We could clearly observe a trend of improved pain scores and functional outcome scores (KOOS, IKDC and Cincinnati scores) with the use of cellular adjuvants (MFx-III-MS, BMAC, PBSC, and SVF). Although the difference in the pain and functional scores improved with the use

of acellular adjuvants (such as PRP, HA, collagen, and AMIC) too in comparison with traditional MFx, the differences were not as substantial as for cellular adjuvants.

This observation is in concurrence with a majority of the studies, which have demonstrated overall improved clinical outcome with acellular (MFx-II) adjuvants. In a prospective, multicenter clinical trial^[31], AMIC with biodegradable type I/III collagen membrane showed significantly improved longer-term radiological (MRI defect filling) and functional outcome (as assessed by Cincinnati and modified ICRS scores) at the 5-year time point, in comparison with MFx-I. In another recent RCT, Shive *et al*^[19] concluded that the use of BST-CarGel (soluble polymer scaffold containing polysaccharide chitosan dispersed in uncoagulated blood) following MFx leads to improved cartilage resurfacing and wound healing. On a similar note, various prospective studies have also reported meliorated outcome (clinical and radiological) following the use of diverse cellular components after MFx (MFx-III). Some such cellular components, which have been successfully tried in cartilage defects, include single-stage cell-based therapy using autologous cartilage fragments (cartilage autograft implantation system-CAIS)^[67], collagen-covered ACI (ACI-C), AMIC^[33], micro-fragmented stromal-vascular fraction (rich in adipose-derived MSCs-ADMSC)^[49], and tri-layered collagen hydroxyapatite biomimetic osteochondral scaffold (CHAS) seeded intra-operatively with autologous chondrocytes (AC) or filtered bone marrow stem/stromal cells (fBMSC)^[99]. In a prospective series by Liu *et al*^[43], it was demonstrated that the application of Kartigen (matrix with autologous bone marrow MSC-derived chondrocyte precursors embedded in atelocollagen) enabled the restoration of columnar surface of articular cartilage, collagen type 2 and glycosaminoglycan in similar composition to native hyaline cartilage (on histology).

Radiological outcome: A majority of the studies reported on MOCART score and MRI filling defect during the follow-up. There was a statistically significant improvement in the MOCART score at the end of 1 year in patients following the use of cellular adjuvants after MFx, indicating a substantially improved cartilage tissue quality and

integration. Although the radiological outcome scores at the subsequent follow-up time points were not statistically different; similar to the clinical outcome, there was a definitive trend towards better outcome after the use of cellular and acellular adjuvants following MFx (cellular > acellular).

In a prospective randomized study by Ibarra *et al*^[59], it was concluded that structural outcome (as assessed by MRI-T2 mapping and MOCART score) and significantly improved clinical outcome (as evaluated by KOOS subscale and Tegner scale) at 1 to 6 years and 4 to 6 years, respectively in patients undergoing matrix-assisted autologous chondrocyte transplantation, as compared with traditional MFx. Patients undergoing adjuvant cell therapy also demonstrated higher response and lower failure rates in this series. Similar prospective cohort studies have demonstrated improved cartilage fill on T2WI MRI and mean MOCART score following surgical treatment with PRP-loaded scaffold (MFx-II)^[100], scaffold augmentation using BMAC (MFx-III)^[100] and transplantation of autologous BMSCs (BMSC-MFx-III)^[60].

Complications and adverse events: Based on our network analysis, we could also clearly identify mitigated complication and failure rates with the higher generations of MFx (although the differences were not statistically significant. In a prospective series by Martincic *et al*^[99], tri-layered CHAS seeded intra-operatively with AC or fBMSC demonstrated significantly improved outcome, in comparison with MFx. In this study, blood soaking of the scaffold prior to cell seeding substantially reduced early post-operative complications like synovitis and arthrofibrosis.

Limitations: Though our ¹ study is one of the most comprehensively-performed reviews of the existing literature on this subject, there are certain limitations. The long-term data on histological and radiological outcomes following recent generations of MFx are limited. There is substantial paucity as well as heterogeneity in the reporting on the diverse functional outcome measures, which prevented uniform comparison of events.

Current status and future directions: Based on our comprehensive review and NMA, we could conclude that the use of acellular and cellular adjuvants (2nd and 3rd generation) marginally improves the overall clinical status (pain and functional scores) and radiological outcome (MOCART score and MRI-filling) in patients undergoing MFx for cartilage defects of the knee. The safety and efficacy of the higher generation MFx procedures are also clearly evident from our review. However, there is a substantial potential for further improvement in the cellular components (chondrocytes over other cellular lineage), culture or processing methodology, delivery modalities (including appropriate scaffolds); as well as better surgical techniques^[6].

CONCLUSION

The use of acellular and cellular adjuvants (2nd and 3rd generation) has shown only marginal improvement in the clinical (pain and functional scores) and radiological outcome (MOCART score and MRI-filling) in patients undergoing MFx for cartilage defects of the knee.

ARTICLE HIGHLIGHTS

Research background

We have noted improvements in the traditional microfracture (MFx) techniques over the decades of its routine use in the management of cartilage defects. The recent generations include the addition of acellular components and cellular components to the cartilage defect. However, the effectiveness of these modifications is not explored further.

Research motivation

To explore the clinical effectiveness of the various generations of the MFx technique to understand their clinical effect in the management of cartilage defects.

Research objectives

To comparatively explore the clinical, radiological and histological outcomes along with the complications reported in the various generations of MFx in the context of the management of cartilage defects.

Research methods

We made a systematic review by utilizing the ⁴ databases such as PubMed, Embase, Web of Science, Cochrane, and Scopus to identify the randomized controlled trials (RCTs) reporting the outcomes of utilization of various generations of MFx in the management of cartilage defects. Network meta-analysis was performed among the three generations for the outcomes analysed using Stata.

Research results

Forty-four RCTs were included in the analysis with patients of mean age of 39.4 (\pm 9.46) years. Upon comparing the results of the other generations with MFx-I as a constant comparator, we noted a trend towards better pain control and functional outcome (KOOS, IKDC and Cincinnati scores) at the end of 1-, 2-, and 5-year time points with MFx-III, although the differences were not statistically significant ($P > 0.05$). We also noted statistically significant MOCART score in the higher generations of MFx (WMD = 17.44; 95%CI: 0.72, 34.16; $P = 0.025$; without significant heterogeneity) at 1 year. However, the difference was not maintained at 2 years. There was a trend towards better defect filling on MRI with the second and third generation MFx, although the difference was not statistically significant ($P > 0.05$).

Research conclusions

The higher generations of traditional MFx technique utilizing acellular and cellular components to augment its potential in the management of cartilage defects has shown only marginal improvement in the clinical and radiological outcomes.

Research perspectives

Future work could focus on the improvement in the cellular components (chondrocytes over other cellular lineage), culture or processing methodology, delivery modalities (including appropriate scaffolds); as well as better surgical techniques to make the clinical impact with their further advancements.

REFERENCES

- 1 **Widuchowski W**, Widuchowski J, Trzaska T. Articular cartilage defects: study of 25,124 knee arthroscopies. *Knee* 2007; **14**: 177-182 [PMID: 17428666 DOI: 10.1016/j.knee.2007.02.001]
- 2 **Farr J**, Cole B, Dhawan A, Kercher J, Sherman S. Clinical cartilage restoration: evolution and overview. *Clin Orthop Relat Res* 2011; **469**: 2696-2705 [PMID: 21240578 DOI: 10.1007/s11999-010-1764-z]
- 3 **Steadman JR**, Rodkey WG, Rodrigo JJ. Microfracture: surgical technique and rehabilitation to treat chondral defects. *Clin Orthop Relat Res* 2001: S362-S369 [PMID: 11603719 DOI: 10.1097/00003086-200110001-00033]
- 4 **Knutsen G**, Engebretsen L, Ludvigsen TC, Drogset JO, Grøntvedt T, Solheim E, Strand T, Roberts S, Isaksen V, Johansen O. Autologous chondrocyte implantation compared with microfracture in the knee. A randomized trial. *J Bone Joint Surg Am* 2004; **86**: 455-464 [PMID: 14996869 DOI: 10.2106/00004623-200403000-00001]
- 5 **Hoemann CD**, Tran-Khanh N, Chevrier A, Chen G, Lascau-Coman V, Mathieu C, Changoor A, Yaroshinsky A, McCormack RG, Stanish WD, Buschmann MD. Chondroinduction Is the Main Cartilage Repair Response to Microfracture and Microfracture With BST-CarGel: Results as Shown by ICRS-II Histological Scoring and a Novel Zonal Collagen Type Scoring Method of Human Clinical Biopsy Specimens. *Am J Sports Med* 2015; **43**: 2469-2480 [PMID: 26260465 DOI: 10.1177/0363546515593943]
- 6 **Muthu S**, Korpershoek JV, Novais EJ, Tawy GF, Hollander AP, Martin I. Failure of cartilage regeneration: emerging hypotheses and related therapeutic strategies. *Nat Rev Rheumatol* 2023; **19**: 403-416 [PMID: 37296196 DOI: 10.1038/s41584-023-00979-5]

- 7 **Bedi A**, Feeley BT, Williams RJ 3rd. Management of articular cartilage defects of the knee. *J Bone Joint Surg Am* 2010; **92**: 994-1009 [PMID: 20360528 DOI: 10.2106/JBJS.I.00895]
- 8 **Krych AJ**, Harnly HW, Rodeo SA, Williams RJ 3rd. Activity levels are higher after osteochondral autograft transfer mosaicplasty than after microfracture for articular cartilage defects of the knee: a retrospective comparative study. *J Bone Joint Surg Am* 2012; **94**: 971-978 [PMID: 22637203 DOI: 10.2106/JBJS.K.00815]
- 9 **Saris DB**, Vanlauwe J, Victor J, Almqvist KF, Verdonk R, Bellemans J, Luyten FP; TIG/ACT/01/2000&EXT Study Group. Treatment of symptomatic cartilage defects of the knee: characterized chondrocyte implantation results in better clinical outcome at 36 months in a randomized trial compared to microfracture. *Am J Sports Med* 2009; **37** Suppl 1: 10S-19S [PMID: 19846694 DOI: 10.1177/0363546509350694]
- 10 **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**: 1986-1996 [PMID: 24051505 DOI: 10.1007/s00167-013-2676-8]
- 11 **Solheim E**, Hegna J, Inderhaug E, Øyen J, Harlem T, Strand T. Results at 10-14 years after microfracture treatment of articular cartilage defects in the knee. *Knee Surg Sports Traumatol Arthrosc* 2016; **24**: 1587-1593 [PMID: 25416965 DOI: 10.1007/s00167-014-3443-1]
- 12 **Case JM**, Scopp JM. Treatment of Articular Cartilage Defects of the Knee With Microfracture and Enhanced Microfracture Techniques. *Sports Med Arthrosc Rev* 2016; **24**: 63-68 [PMID: 27135288 DOI: 10.1097/JSA.0000000000000113]
- 13 **Koh YG**, Kwon OR, Kim YS, Choi YJ, Tak DH. Adipose-Derived Mesenchymal Stem Cells With Microfracture Versus Microfracture Alone: 2-Year Follow-up of a Prospective Randomized Trial. *Arthroscopy* 2016; **32**: 97-109 [PMID: 26585585 DOI: 10.1016/j.arthro.2015.09.010]
- 14 **Lee KB**, Wang VT, Chan YH, Hui JH. A novel, minimally-invasive technique of cartilage repair in the human knee using arthroscopic microfracture and injections of

mesenchymal stem cells and hyaluronic acid--a prospective comparative study on safety and short-term efficacy. *Ann Acad Med Singap* 2012; **41**: 511-517 [PMID: 23235728]

15 **Lee GW**, Son JH, Kim JD, Jung GH. Is platelet-rich plasma able to enhance the results of arthroscopic microfracture in early osteoarthritis and cartilage lesion over 40 years of age? *Eur J Orthop Surg Traumatol* 2013; **23**: 581-587 [PMID: 23412171 DOI: 10.1007/s00590-012-1038-4]

16 **Siclari A**, Mascaro G, Gentili C, Kaps C, Cancedda R, Boux E. Cartilage repair in the knee with subchondral drilling augmented with a platelet-rich plasma-immersed polymer-based implant. *Knee Surg Sports Traumatol Arthrosc* 2014; **22**: 1225-1234 [PMID: 23563814 DOI: 10.1007/s00167-013-2484-1]

17 **Saw KY**, Hussin P, Loke SC, Azam M, Chen HC, Tay YG, Low S, Wallin KL, Ragavanaidu K. Articular cartilage regeneration with autologous marrow aspirate and hyaluronic Acid: an experimental study in a goat model. *Arthroscopy* 2009; **25**: 1391-1400 [PMID: 19962065 DOI: 10.1016/j.arthro.2009.07.011]

18 **Strauss E**, Schachter A, Frenkel S, Rosen J. The efficacy of intra-articular hyaluronan injection after the microfracture technique for the treatment of articular cartilage lesions. *Am J Sports Med* 2009; **37**: 720-726 [PMID: 19204370 DOI: 10.1177/0363546508328415]

19 **Shive MS**, Stanish WD, McCormack R, Forriol F, Mohtadi N, Pelet S, Desnoyers J, Méthot S, Vehik K, Restrepo A. BST-CarGel® Treatment Maintains Cartilage Repair Superiority over Microfracture at 5 Years in a Multicenter Randomized Controlled Trial. *Cartilage* 2015; **6**: 62-72 [PMID: 26069709 DOI: 10.1177/1947603514562064]

20 **Stanish WD**, McCormack R, Forriol F, Mohtadi N, Pelet S, Desnoyers J, Restrepo A, Shive MS. Novel scaffold-based BST-CarGel treatment results in superior cartilage repair compared with microfracture in a randomized controlled trial. *J Bone Joint Surg Am* 2013; **95**: 1640-1650 [PMID: 24048551 DOI: 10.2106/jbjs.l.01345]

21 **Strauss EJ**, Barker JU, Kercher JS, Cole BJ, Mithoefer K. Augmentation Strategies following the Microfracture Technique for Repair of Focal Chondral Defects. *Cartilage* 2010; **1**: 145-152 [PMID: 26069546 DOI: 10.1177/1947603510366718]

- 22 **Hutton B**, Salanti G, Caldwell DM, Chaimani A, Schmid CH, Cameron C, Ioannidis JP, Straus S, Thorlund K, Jansen JP, Mulrow C, Catalá-López F, Gøtzsche PC, Dickersin K, Boutron I, Altman DG, Moher D. The PRISMA extension statement for reporting of systematic reviews incorporating network meta-analyses of health care interventions: checklist and explanations. *Ann Intern Med* 2015; **162**: 777-784 [PMID: 26030634 DOI: 10.7326/M14-2385]
- 23 **McGowan J**, Sampson M, Salzwedel DM, Cogo E, Foerster V, Lefebvre C. PRESS Peer Review of Electronic Search Strategies: 2015 Guideline Statement. *J Clin Epidemiol* 2016; **75**: 40-46 [PMID: 27005575 DOI: 10.1016/j.jclinepi.2016.01.021]
- 24 **Orthopaedic Research Group**. ORG Cartilage Treatment Classifier Tool. [cited 12 December 2023]. Available from: <https://orthopaedicresearchgroup.com/contact.php>
- 25 **Sterne JAC**, Savović J, Page MJ, Elbers RG, Blencowe NS, Boutron I, Cates CJ, Cheng HY, Corbett MS, Eldridge SM, Emberson JR, Hernán MA, Hopewell S, Hróbjartsson A, Junqueira DR, Jüni P, Kirkham JJ, Lasserson T, Li T, McAleenan A, Reeves BC, Shepperd S, Shrier I, Stewart LA, Tilling K, White IR, Whiting PF, Higgins JPT. RoB 2: a revised tool for assessing risk of bias in randomised trials. *BMJ* 2019; **366**: 14898 [PMID: 31462531 DOI: 10.1136/bmj.14898]
- 26 **Hwang H**, DeSantis SM. Multivariate network meta-analysis to mitigate the effects of outcome reporting bias. *Stat Med* 2018; **37**: 3254-3266 [PMID: 29882392 DOI: 10.1002/sim.7815]
- 27 **White IR**. Network Meta-analysis. *Stata J* 2015; **15**: 951-985 [DOI: 10.1177/1536867X1501500403]
- 28 **Lu G**, Ades A. Modeling between-trial variance structure in mixed treatment comparisons. *Biostatistics* 2009; **10**: 792-805 [PMID: 19687150 DOI: 10.1093/biostatistics/kxp032]
- 29 **Nikolakopoulou A**, Higgins JPT, Papakonstantinou T, Chaimani A, Del Giovane C, Egger M, Salanti G. CINeMA: An approach for assessing confidence in the results of a network meta-analysis. *PLoS Med* 2020; **17**: e1003082 [PMID: 32243458 DOI: 10.1371/journal.pmed.1003082]

- 30 **Papakonstantinou T**, Nikolakopoulou A, Higgins JPT, Egger M, Salanti G. CINeMA: Software for semiautomated assessment of the confidence in the results of network meta-analysis. *Campbell Syst Rev* 2020; **16**: e1080 [PMID: 37131978 DOI: 10.1002/cl2.1080]
- 31 **Volz M**, Schaumburger J, Frick H, Grifka J, Anders S. A randomized controlled trial demonstrating sustained benefit of Autologous Matrix-Induced Chondrogenesis over microfracture at five years. *Int Orthop* 2017; **41**: 797-804 [PMID: 28108777 DOI: 10.1007/s00264-016-3391-0]
- 32 **Niemeyer P**, Laute V, Zinser W, Becher C, Kolombe T, Fay J, Pietsch S, Kuźma T, Widuchowski W, Fickert S. A Prospective, Randomized, Open-Label, Multicenter, Phase III Noninferiority Trial to Compare the Clinical Efficacy of Matrix-Associated Autologous Chondrocyte Implantation With Spheroid Technology Versus Arthroscopic Microfracture for Cartilage Defects of the Knee. *Orthop J Sports Med* 2019; **7**: 2325967119854442 [PMID: 31317047 DOI: 10.1177/2325967119854442]
- 33 **Fossum V**, Hansen AK, Wilsgaard T, Knutsen G. Collagen-Covered Autologous Chondrocyte Implantation Versus Autologous Matrix-Induced Chondrogenesis: A Randomized Trial Comparing 2 Methods for Repair of Cartilage Defects of the Knee. *Orthop J Sports Med* 2019; **7**: 2325967119868212 [PMID: 31555714 DOI: 10.1177/2325967119868212]
- 34 **Ulstein S**, Årøen A, Røtterud JH, Løken S, Engebretsen L, Heir S. Microfracture technique versus osteochondral autologous transplantation mosaicplasty in patients with articular chondral lesions of the knee: a prospective randomized trial with long-term follow-up. *Knee Surg Sports Traumatol Arthrosc* 2014; **22**: 1207-1215 [PMID: 24441734 DOI: 10.1007/s00167-014-2843-6]
- 35 **Visna P**, Pasa L, Cizmár I, Hart R, Hoch J. Treatment of deep cartilage defects of the knee using autologous chondrograft transplantation and by abrasive techniques--a randomized controlled study. *Acta Chir Belg* 2004; **104**: 709-714 [PMID: 15663280 DOI: 10.1080/00015458.2004.11679648]

- 36 **Van Assche D**, Staes F, Van Caspel D, Vanlauwe J, Bellemans J, Saris DB, Luyten FP. Autologous chondrocyte implantation versus microfracture for knee cartilage injury: a prospective randomized trial, with 2-year follow-up. *Knee Surg Sports Traumatol Arthrosc* 2010; **18**: 486-495 [PMID: 19820916 DOI: 10.1007/s00167-009-0955-1]
- 37 **Saw KY**, Anz A, Siew-Yoke Jee C, Merican S, Ching-Soong Ng R, Roohi SA, Ragavanaidu K. Articular cartilage regeneration with autologous peripheral blood stem cells versus hyaluronic acid: a randomized controlled trial. *Arthroscopy* 2013; **29**: 684-694 [PMID: 23380230 DOI: 10.1016/j.arthro.2012.12.008]
- 38 **Anders S**, Volz M, Frick H, Gellissen J. A Randomized, Controlled Trial Comparing Autologous Matrix-Induced Chondrogenesis (AMIC®) to Microfracture: Analysis of 1- and 2-Year Follow-Up Data of 2 Centers. *Open Orthop J* 2013; **7**: 133-143 [PMID: 23730377 DOI: 10.2174/1874325001307010133]
- 39 **Brittberg M**, Recker D, Ilgenfritz J, Saris DBF; SUMMIT Extension Study Group. Matrix-Applied Characterized Autologous Cultured Chondrocytes Versus Microfracture: Five-Year Follow-up of a Prospective Randomized Trial. *Am J Sports Med* 2018; **46**: 1343-1351 [PMID: 29565642 DOI: 10.1177/0363546518756976]
- 40 **Lim HC**, Bae JH, Song SH, Park YE, Kim SJ. Current treatments of isolated articular cartilage lesions of the knee achieve similar outcomes. *Clin Orthop Relat Res* 2012; **470**: 2261-2267 [PMID: 22422593 DOI: 10.1007/s11999-012-2304-9]
- 41 **Knutsen G**, Drogset JO, Engebretsen L, Grøntvedt T, Isaksen V, Ludvigsen TC, Roberts S, Solheim E, Strand T, Johansen O. A randomized trial comparing autologous chondrocyte implantation with microfracture. Findings at five years. *J Bone Joint Surg Am* 2007; **89**: 2105-2112 [PMID: 17908884 DOI: 10.2106/JBJS.G.00003]
- 42 **Knutsen G**, Drogset JO, Engebretsen L, Grøntvedt T, Ludvigsen TC, Løken S, Solheim E, Strand T, Johansen O. A Randomized Multicenter Trial Comparing Autologous Chondrocyte Implantation with Microfracture: Long-Term Follow-up at 14 to 15 Years. *J Bone Joint Surg Am* 2016; **98**: 1332-1339 [PMID: 27535435 DOI: 10.2106/JBJS.15.01208]

- 43 **Liu YL**, Yen CC, Liu TT, Chang CH, Shih TT, Wang JH, Yang MC, Lin FH, Liu HC. Safety and Efficacy of Kartigen(®) in Treating Cartilage Defects: A Randomized, Controlled, Phase I Trial. *Polymers (Basel)* 2021; **13** [PMID: 34577930 DOI: 10.3390/polym13183029]
- 44 **Yoon KH**, Park JY, Lee JY, Lee E, Lee J, Kim SG. Costal Chondrocyte-Derived Pellet-Type Autologous Chondrocyte Implantation for Treatment of Articular Cartilage Defect. *Am J Sports Med* 2020; **48**: 1236-1245 [PMID: 32125878 DOI: 10.1177/0363546520905565]
- 45 **Kon E**, Filardo G, Brittberg M, Busacca M, Condello V, Engebretsen L, Marlovits S, Niemeyer P, Platzer P, Posthumus M, Verdonk P, Verdonk R, Victor J, van der Merwe W, Widuchowski W, Zorzi C, Marcacci M. A multilayer biomaterial for osteochondral regeneration shows superiority vs microfractures for the treatment of osteochondral lesions in a multicentre randomized trial at 2 years. *Knee Surg Sports Traumatol Arthrosc* 2018; **26**: 2704-2715 [PMID: 28913600 DOI: 10.1007/s00167-017-4707-3]
- 46 **Vanlauwe J**, Saris DB, Victor J, Almqvist KF, Bellemans J, Luyten FP; TIG/ACT/01/2000&EXT Study Group. Five-year outcome of characterized chondrocyte implantation versus microfracture for symptomatic cartilage defects of the knee: early treatment matters. *Am J Sports Med* 2011; **39**: 2566-2574 [PMID: 21908720 DOI: 10.1177/0363546511422220]
- 47 **Basad E**, Ishaque B, Bachmann G, Stürz H, Steinmeyer J. Matrix-induced autologous chondrocyte implantation versus microfracture in the treatment of cartilage defects of the knee: a 2-year randomised study. *Knee Surg Sports Traumatol Arthrosc* 2010; **18**: 519-527 [PMID: 20062969 DOI: 10.1007/s00167-009-1028-1]
- 48 **Solheim E**, Hegna J, Strand T, Harlem T, Inderhaug E. Randomized Study of Long-term (15-17 Years) Outcome After Microfracture Versus Mosaicplasty in Knee Articular Cartilage Defects. *Am J Sports Med* 2018; **46**: 826-831 [PMID: 29253350 DOI: 10.1177/0363546517745281]
- 49 **Bisicchia S**, Bernardi G, Pagnotta SM, Tudisco C. Micro-fragmented stromal-vascular fraction plus microfractures provides better clinical results than microfractures

alone in symptomatic focal chondral lesions of the knee. *Knee Surg Sports Traumatol Arthrosc* 2020; **28**: 1876-1884 [PMID: 31297576 DOI: 10.1007/s00167-019-05621-0]

50 **Saris D**, Price A, Widuchowski W, Bertrand-Marchand M, Caron J, Drogset JO, Emans P, Podskubka A, Tsuchida A, Kili S, Levine D, Brittberg M; SUMMIT study group. Matrix-Applied Characterized Autologous Cultured Chondrocytes Versus Microfracture: Two-Year Follow-up of a Prospective Randomized Trial. *Am J Sports Med* 2014; **42**: 1384-1394 [PMID: 24714783 DOI: 10.1177/0363546514528093]

51 **Saris DB**, Vanlauwe J, Victor J, Haspl M, Bohnsack M, Fortems Y, Vandekerckhove B, Almqvist KF, Claes T, Handelberg F, Lagae K, van der Bauwhede J, Vandenuecker H, Yang KG, Jelic M, Verdonk R, Veulemans N, Bellemans J, Luyten FP. Characterized chondrocyte implantation results in better structural repair when treating symptomatic cartilage defects of the knee in a randomized controlled trial versus microfracture. *Am J Sports Med* 2008; **36**: 235-246 [PMID: 18202295 DOI: 10.1177/0363546507311095]

52 **Qiao Z**, Tang J, Yue B, Wang J, Zhang J, Xuan L, Dai C, Li S, Li M, Xu C, Dai K, Wang Y. Human adipose-derived mesenchymal progenitor cells plus microfracture and hyaluronic acid for cartilage repair: a Phase IIa trial. *Regen Med* 2020; **15**: 1193-1214 [PMID: 32043426 DOI: 10.2217/rme-2019-0068]

53 **Nguyen PD**, Tran TD, Nguyen HT, Vu HT, Le PT, Phan NL, Vu NB, Phan NK, Van Pham P. Comparative Clinical Observation of Arthroscopic Microfracture in the Presence and Absence of a Stromal Vascular Fraction Injection for Osteoarthritis. *Stem Cells Transl Med* 2017; **6**: 187-195 [PMID: 28170179 DOI: 10.5966/sctm.2016-0023]

54 **Lim HC**, Park YB, Ha CW, Cole BJ, Lee BK, Jeong HJ, Kim MK, Bin SI, Choi CH, Choi CH, Yoo JD; Cartistem Research Group, Yoon JR, Chung JY. Allogeneic Umbilical Cord Blood-Derived Mesenchymal Stem Cell Implantation Versus Microfracture for Large, Full-Thickness Cartilage Defects in Older Patients: A Multicenter Randomized Clinical Trial and Extended 5-Year Clinical Follow-up. *Orthop J Sports Med* 2021; **9**: 2325967120973052 [PMID: 33490296 DOI: 10.1177/2325967120973052]

55 **Venosa M**, Calafiore F, Mazzoleni M, Romanini E, Cerciello S, Calvisi V. Platelet-Rich Plasma and Adipose-Derived Mesenchymal Stem Cells in Association with

Arthroscopic Microfracture of Knee Articular Cartilage Defects: A Pilot Randomized Controlled Trial. *Adv Orthop* 2022; **2022**: 6048477 [PMID: 35529427 DOI: 10.1155/2022/6048477]

56 **Kim MS**, Koh IJ, Choi YJ, Pak KH, In Y. Collagen Augmentation Improves the Quality of Cartilage Repair After Microfracture in Patients Undergoing High Tibial Osteotomy: A Randomized Controlled Trial. *Am J Sports Med* 2017; **45**: 1845-1855 [PMID: 28282221 DOI: 10.1177/0363546517691942]

57 **Kim MS**, Chun CH, Wang JH, Kim JG, Kang SB, Yoo JD, Chon JG, Kim MK, Moon CW, Chang CB, Song IS, Ha JK, Choi NY, In Y. Microfractures Versus a Porcine-Derived Collagen-Augmented Chondrogenesis Technique for Treating Knee Cartilage Defects: A Multicenter Randomized Controlled Trial. *Arthroscopy* 2020; **36**: 1612-1624 [PMID: 31785390 DOI: 10.1016/j.arthro.2019.11.110]

58 **Kane MS**, Williams RJ III, DeBerardino TM, Taylor D, Ma CB, Anderson DE, Crawford DC. Review of an exploratory phase II FDA regulated clinical trial of a novel surgical innovation: completion of a prospective, randomized, controlled trial to compare NeoCart with the standard-of-care, microfracture, for articular cartilage repair. *Annals of Joint* 2018; **3** [DOI: 10.21037/aoj.2018.06.08]

59 **Ibarra C**, Villalobos E, Madrazo-Ibarra A, Velasquillo C, Martinez-Lopez V, Izaguirre A, Olivos-Meza A, Cortes-Gonzalez S, Perez-Jimenez FJ, Vargas-Ramirez A, Franco-Sanchez G, Ibarra-Ibarra LG, Sierra-Suarez L, Almazan A, Ortega-Sanchez C, Trueba C, Martin FB, Arredondo-Valdes R, Chavez-Arias D. Arthroscopic Matrix-Assisted Autologous Chondrocyte Transplantation Versus Microfracture: A 6-Year Follow-up of a Prospective Randomized Trial. *Am J Sports Med* 2021; **49**: 2165-2176 [PMID: 34048286 DOI: 10.1177/03635465211010487]

60 **Hashimoto Y**, Nishida Y, Takahashi S, Nakamura H, Mera H, Kashiwa K, Yoshiya S, Inagaki Y, Uematsu K, Tanaka Y, Asada S, Akagi M, Fukuda K, Hosokawa Y, Myoui A, Kamei N, Ishikawa M, Adachi N, Ochi M, Wakitani S. Transplantation of autologous bone marrow-derived mesenchymal stem cells under arthroscopic surgery with microfracture versus microfracture alone for articular cartilage lesions in the knee: A

multicenter prospective randomized control clinical trial. *Regen Ther* 2019; **11**: 106-113 [PMID: 31312692 DOI: 10.1016/j.reth.2019.06.002]

61 **Gudas R**, Stankevicius E, Monastyreckiene E, Pranys D, Kalesinskas RJ. Osteochondral autologous transplantation versus microfracture for the treatment of articular cartilage defects in the knee joint in athletes. *Knee Surg Sports Traumatol Arthrosc* 2006; **14**: 834-842 [PMID: 16552548 DOI: 10.1007/s00167-006-0067-0]

62 **Gudas R**, Gudaitė A, Mickevičius T, Masiulis N, Simonaitytė R, Cekanauskas E, Skurvydas A. Comparison of osteochondral autologous transplantation, microfracture, or debridement techniques in articular cartilage lesions associated with anterior cruciate ligament injury: a prospective study with a 3-year follow-up. *Arthroscopy* 2013; **29**: 89-97 [PMID: 23142295 DOI: 10.1016/j.arthro.2012.06.009]

63 **Gudas R**, Kalesinskas RJ, Kimtys V, Stankevicius E, Toliulis V, Bernotavicius G, Smailys A. A prospective randomized clinical study of mosaic osteochondral autologous transplantation versus microfracture for the treatment of osteochondral defects in the knee joint in young athletes. *Arthroscopy* 2005; **21**: 1066-1075 [PMID: 16171631 DOI: 10.1016/j.arthro.2005.06.018]

64 **Glasbrenner J**, Petersen W, Raschke MJ, Steiger M, Verdonk R, Castelli CC, Zappalà G, Fritschy D, Herbort M. Matrix-Augmented Bone Marrow Stimulation With a Polyglycolic Acid Membrane With Hyaluronan vs Microfracture in Local Cartilage Defects of the Femoral Condyles: A Multicenter Randomized Controlled Trial. *Orthop J Sports Med* 2020; **8**: 2325967120922938 [PMID: 32528994 DOI: 10.1177/2325967120922938]

65 **Dasar U**, Gursay S, Akkaya M, Algin O, Isik C, Bozkurt M. Microfracture technique versus carbon fibre rod implantation for treatment of knee articular cartilage lesions. *J Orthop Surg (Hong Kong)* 2016; **24**: 188-193 [PMID: 27574261 DOI: 10.1177/1602400214]

66 **Crawford DC**, DeBerardino TM, Williams RJ 3rd. NeoCart, an autologous cartilage tissue implant, compared with microfracture for treatment of distal femoral cartilage lesions: an FDA phase-II prospective, randomized clinical trial after two years. *J Bone Joint Surg Am* 2012; **94**: 979-989 [PMID: 22637204 DOI: 10.2106/JBJS.K.00533]

- 67 **Cole BJ**, Farr J, Winalski CS, Hosea T, Richmond J, Mandelbaum B, De Deyne PG. Outcomes after a single-stage procedure for cell-based cartilage repair: a prospective clinical safety trial with 2-year follow-up. *Am J Sports Med* 2011; **39**: 1170-1179 [PMID: 21460066 DOI: 10.1177/0363546511399382]
- 68 **Chung JY**, Lee DH, Kim TH, Kwack KS, Yoon KH, Min BH. Cartilage extra-cellular matrix biomembrane for the enhancement of microfractured defects. *Knee Surg Sports Traumatol Arthrosc* 2014; **22**: 1249-1259 [PMID: 24258020 DOI: 10.1007/s00167-013-2716-4]
- 69 **Barié A**, Kruck P, Sorbi R, Rehnitz C, Oberle D, Walker T, Zeifang F, Moradi B. Prospective Long-term Follow-up of Autologous Chondrocyte Implantation With Periosteum Versus Matrix-Associated Autologous Chondrocyte Implantation: A Randomized Clinical Trial. *Am J Sports Med* 2020; **48**: 2230-2241 [PMID: 32667270 DOI: 10.1177/0363546520928337]
- 70 **Clavé A**, Potel JF, Servien E, Neyret P, Dubrana F, Stindel E. Third-generation autologous chondrocyte implantation versus mosaicplasty for knee cartilage injury: 2-year randomized trial. *J Orthop Res* 2016; **34**: 658-665 [PMID: 26742454 DOI: 10.1002/jor.23152]
- 71 **de Queiroz AAB**, Debieux P, Amaro J, Ferretti M, Cohen M. Hydrogel implant is as effective as osteochondral autologous transplantation for treating focal cartilage knee injury in 24 months. *Knee Surg Sports Traumatol Arthrosc* 2018; **26**: 2934-2941 [PMID: 29335748 DOI: 10.1007/s00167-018-4834-5]
- 72 **Gooding CR**, Bartlett W, Bentley G, Skinner JA, Carrington R, Flanagan A. A prospective, randomised study comparing two techniques of autologous chondrocyte implantation for osteochondral defects in the knee: Periosteum covered versus type I/III collagen covered. *Knee* 2006; **13**: 203-210 [PMID: 16644224 DOI: 10.1016/j.knee.2006.02.011]
- 73 **Bartlett W**, Skinner JA, Gooding CR, Carrington RW, Flanagan AM, Briggs TW, Bentley G. Autologous chondrocyte implantation versus matrix-induced autologous chondrocyte implantation for osteochondral defects of the knee: a prospective,

randomised study. *J Bone Joint Surg Br* 2005; **87**: 640-645 [PMID: 15855365 DOI: 10.1302/0301-620X.87B5.15905]

74 **Bentley G**, Biant LC, Carrington RW, Akmal M, Goldberg A, Williams AM, Skinner JA, Pringle J. A prospective, randomised comparison of autologous chondrocyte implantation versus mosaicplasty for osteochondral defects in the knee. *J Bone Joint Surg Br* 2003; **85**: 223-230 [PMID: 12678357 DOI: 10.1302/0301-620x.85b2.13543]

75 **Zeifang F**, Oberle D, Nierhoff C, Richter W, Moradi B, Schmitt H. Autologous chondrocyte implantation using the original periosteum-cover technique versus matrix-associated autologous chondrocyte implantation: a randomized clinical trial. *Am J Sports Med* 2010; **38**: 924-933 [PMID: 19966102 DOI: 10.1177/0363546509351499]

76 **Bentley G**, Biant LC, Vijayan S, Macmull S, Skinner JA, Carrington RW. Minimum ten-year results of a prospective randomised study of autologous chondrocyte implantation versus mosaicplasty for symptomatic articular cartilage lesions of the knee. *J Bone Joint Surg Br* 2012; **94**: 504-509 [PMID: 22434467 DOI: 10.1302/0301-620X.94B4.27495]

77 **Zaffagnini S**, Boffa A, Andriolo L, Reale D, Busacca M, Di Martino A, Filardo G. Mosaicplasty vs Matrix-Assisted Autologous Chondrocyte Transplantation for Knee Cartilage Defects: A Long-Term Clinical and Imaging Evaluation. *App Sci-Basel* 2020; **10**: 4615 [DOI: 10.3390/app10134615]

78 **GBD 2017 Disease and Injury Incidence and Prevalence Collaborators**. Global, regional, and national incidence, prevalence, and years lived with disability for 354 diseases and injuries for 195 countries and territories, 1990-2017: a systematic analysis for the Global Burden of Disease Study 2017. *Lancet* 2018; **392**: 1789-1858 [PMID: 30496104 DOI: 10.1016/S0140-6736(18)32279-7]

79 **Curl WW**, Krome J, Gordon ES, Rushing J, Smith BP, Poehling GG. Cartilage injuries: a review of 31,516 knee arthroscopies. *Arthroscopy* 1997; **13**: 456-460 [PMID: 9276052 DOI: 10.1016/s0749-8063(97)90124-9]

- 80 **Hjelle K**, Solheim E, Strand T, Muri R, Brittberg M. Articular cartilage defects in 1,000 knee arthroscopies. *Arthroscopy* 2002; **18**: 730-734 [PMID: 12209430 DOI: 10.1053/jars.2002.32839]
- 81 **Arøen A**, Løken S, Heir S, Alvik E, Ekeland A, Granlund OG, Engebretsen L. Articular cartilage lesions in 993 consecutive knee arthroscopies. *Am J Sports Med* 2004; **32**: 211-215 [PMID: 14754746 DOI: 10.1177/0363546503259345]
- 82 **Hunter W**. Of the structure and disease of articulating cartilages. 1743. *Clin Orthop Relat Res* 1995: 3-6 [PMID: 7671493]
- 83 **William H**. Of the structure and diseases of articulating cartilages. *Phil. Trans R Soc* 1743; **42**: 514-521 [DOI: 10.1098/rstl.1742.0079]
- 84 **Buckwalter JA**. Articular cartilage: injuries and potential for healing. *J Orthop Sports Phys Ther* 1998; **28**: 192-202 [PMID: 9785255 DOI: 10.2519/jospt.1998.28.4.192]
- 85 **Muthu S**. Osteoarthritis, an old wine in a new bottle!. *World J Orthop* 2023; **14**: 1-5 [PMID: 36686283 DOI: 10.5312/wjo.v14.i1.1]
- 86 **Kim HK**, Moran ME, Salter RB. The potential for regeneration of articular cartilage in defects created by chondral shaving and subchondral abrasion. An experimental investigation in rabbits. *J Bone Joint Surg Am* 1991; **73**: 1301-1315 [PMID: 1918112]
- 87 **Shapiro F**, Koide S, Glimcher MJ. Cell origin and differentiation in the repair of full-thickness defects of articular cartilage. *J Bone Joint Surg Am* 1993; **75**: 532-553 [PMID: 8478382 DOI: 10.2106/00004623-199304000-00009]
- 88 **Everhart JS**, Campbell AB, Abouljoud MM, Kirven JC, Flanigan DC. Cost-efficacy of Knee Cartilage Defect Treatments in the United States. *Am J Sports Med* 2020; **48**: 242-251 [PMID: 31038980 DOI: 10.1177/0363546519834557]
- 89 **Hunziker EB**, Lippuner K, Keel MJ, Shintani N. An educational review of cartilage repair: precepts & practice--myths & misconceptions--progress & prospects. *Osteoarthritis Cartilage* 2015; **23**: 334-350 [PMID: 25534362 DOI: 10.1016/j.joca.2014.12.011]
- 90 **Niemeyer P**, Becher C, Brucker PU, Buhs M, Fickert S, Gelse K, Günther D, Kaelin R, Kreuz P, Lützner J, Nehrer S, Madry H, Marlovits S, Mehl J, Ott H, Pietschmann M,

Spahn G, Tischer T, Volz M, Walther M, Welsch G, Zellner J, Zinser W, Angele P. [Significance of Matrix-augmented Bone Marrow Stimulation for Treatment of Cartilage Defects of the Knee: A Consensus Statement of the DGOU Working Group on Tissue Regeneration]. *Z Orthop Unfall* 2018; **156**: 513-532 [PMID: 29913540 DOI: 10.1055/a-0591-6457]

91 **Zheng MH**, Willers C, Kirilak L, Yates P, Xu J, Wood D, Shimmin A. Matrix-induced autologous chondrocyte implantation (MACI): biological and histological assessment. *Tissue Eng* 2007; **13**: 737-746 [PMID: 17371156 DOI: 10.1089/ten.2006.0246]

92 **Wang CC**, Yang KC, Lin KH, Liu YL, Liu HC, Lin FH. Cartilage regeneration in SCID mice using a highly organized three-dimensional alginate scaffold. *Biomaterials* 2012; **33**: 120-127 [PMID: 21982587 DOI: 10.1016/j.biomaterials.2011.09.042]

93 **Vacanti JP**. Beyond transplantation. Third annual Samuel Jason Mixter lecture. *Arch Surg* 1988; **123**: 545-549 [PMID: 3282491 DOI: 10.1001/archsurg.1988.01400290027003]

94 **Brittberg M**, Lindahl A, Nilsson A, Ohlsson C, Isaksson O, Peterson L. Treatment of deep cartilage defects in the knee with autologous chondrocyte transplantation. *N Engl J Med* 1994; **331**: 889-895 [PMID: 8078550 DOI: 10.1056/NEJM199410063311401]

95 **Teo AQA**, Wong KL, Shen L, Lim JY, Toh WS, Lee EH, Hui JHP. Equivalent 10-Year Outcomes After Implantation of Autologous Bone Marrow-Derived Mesenchymal Stem Cells Versus Autologous Chondrocyte Implantation for Chondral Defects of the Knee. *Am J Sports Med* 2019; **47**: 2881-2887 [PMID: 31433674 DOI: 10.1177/0363546519867933]

96 **Pittenger MF**, Mackay AM, Beck SC, Jaiswal RK, Douglas R, Mosca JD, Moorman MA, Simonetti DW, Craig S, Marshak DR. Multilineage potential of adult human mesenchymal stem cells. *Science* 1999; **284**: 143-147 [PMID: 10102814 DOI: 10.1126/science.284.5411.143]

97 **Prajwal GS**, Jeyaraman N, Kanth V K, Jeyaraman M, Muthu S, Rajendran SNS, Rajendran RL, Khanna M, Oh EJ, Choi KY, Chung HY, Ahn BC, Gangadaran P. Lineage Differentiation Potential of Different Sources of Mesenchymal Stem Cells for Osteoarthritis Knee. *Pharmaceuticals (Basel)* 2022; **15** [PMID: 35455383 DOI: 10.3390/ph15040386]

98 **Devitt BM**, Bell SW, Webster KE, Feller JA, Whitehead TS. Surgical treatments of cartilage defects of the knee: Systematic review of randomised controlled trials. *Knee* 2017; **24**: 508-517 [PMID: 28189406 DOI: 10.1016/j.knee.2016.12.002]

99 **Martinčič D**, Leban J, Filardo G, Busacca M, Barlič A, Veber M, Drobnič M. Autologous chondrocytes versus filtered bone marrow mesenchymal stem/stromal cells for knee cartilage repair-a prospective study. *Int Orthop* 2021; **45**: 931-939 [PMID: 32712785 DOI: 10.1007/s00264-020-04727-2]

100 **Krych AJ**, Nawabi DH, Farshad-Amacker NA, Jones KJ, Maak TG, Potter HG, Williams RJ 3rd. 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**: 91-98 [PMID: 26574602 DOI:

Figure 1 Preferred Reporting Items for Systematic Review and Meta-analysis flow diagram of selection of studies included in the analysis. RCT: Randomized controlled trial.

Figure 2 Forest plot comparing the generations of microfracture for the functional outcomes reported at 1 year among the included studies in the network. 95%CI: 95% confidence interval; VAS: Visual Analog Scale; MFx: Microfracture; KOOS: Knee Osteoarthritis Outcome Scale; IKDC: International Knee Documentation Committee;

Figure 3 Forest plot comparing the generations of microfracture for the radiological outcomes reported at 1 year among the included studies in the network. 95%CI: 95% confidence interval; MFx: Microfracture; MRI: Magnetic resonance imaging; MOCART: Magnetic resonance observation of cartilage repair tissue.

Figure 4 Forest plot comparing the generations of microfracture for the functional and radiological outcomes at 2 years reported among the included studies in the

network. 95%CI: 95% confidence interval; VAS: Visual Analog Scale; MFx: Microfracture; KOOS: Knee Osteoarthritis Outcome Scale; IKDC: International Knee Documentation Committee; MRI: Magnetic resonance imaging; MOCART: Magnetic resonance observation of cartilage repair tissue.

Figure 5 Forest plot comparing the generations of microfracture for the functional outcomes reported at 5 years among the included studies in the network. 95%CI: 95% confidence interval; mFX: Microfracture; IKDC: International Knee Documentation Committee.

Table 1 Characteristics of included studies in the network meta-analysis, each row depicts the individual comparator arm in the studies included

Study ID	Author	Year	Country	Study design	Sample size		Treatment		Mean age		Female		Mean defect size		Follow-up (months)
					Control	Intervention	Control	Intervention	Control	Intervention	Control	Intervention	Control	Intervention	
1	A Volz	2017	Germany	RCT	34	13	AMIC	Microfracture	40.0	36.5	7	3	3.9	2.9	60
2	P Niemeyer	2019	Germany	RCT	52	50	MACI	Microfracture	36.0	37.0	19	22	2.7	2.4	24
3	V Fossum	2019	Norway	RCT	21	20	ACI-C	AMIC	37.2	38.3	7	12	4.9	5.2	24
4	S Ulstein	2014	Norway	RCT	11	14	Microfracture	AOT	31.7	32.7	11	9	2.6	3.0	120
5	P Visna	2004	Czech Republic	RCT	25	25	Autologous chondrograft transplantation	Microfracture	29.4	32.2	7	9	4	3.3	12
6	DV Assche	2010	Belgium	RCT	33	34	ACI-P	Microfracture	34.0	34.0	11	10	2.5	2.3	24
7	K Saw	2013	United States	RCT	24	25	Microfracture with HA	Microfracture with PBSC	42.0	38.0	17	15	NA	NA	18
8	S Anders	2013	Germany	RCT	22	8	AMIC	Microfracture	41.0	38.0	17	15	3.7	3.5	24
9	GW Lee	2013	Republic of Korea	RCT	25	24	Microfracture	Microfracture with PRP	46.0	46.0	10	10	3.0	3.0	24
10	M Brittberg	2018	Sweden	RCT	65	63	MACI	Microfracture	38.0	34.0	23	20	5.1	4.9	60
11	H Lim	2012	South Korea	RCT	30	22	Microfracture	AOT	32.9	30.4	12	10	2.7	2.7	60
12	G Knutson	2007	Norway	RCT	40	18	ACI-P	ACI-P	25.1	25.1		8		2.8	60
13	G Knutson	2016	Norway	RCT	40	40	ACI-P	Microfracture	33.3	31.1			5.0	5.0	60
14	Y Liu	2021	Taiwan	RCT	10	5	Kartigen	Microfracture	54.8	67.8	5	3	2.9	1.0	180
15	K Yoon	2020	Republic of Korea	RCT	20	10	ACI-CCP	Microfracture	41.5	47.2	6	7	3.5	2.5	12
16	E Kon	2017	Italy	RCT	51	49	Collagen HA	Microfracture	34.0	35.2	15	18	3.4	3.4	24
17	J Vanlauwe	2011	Belgium	RCT	51	61	ACI-P	Microfracture	33.9	33.9	22	20	2.6	2.4	60
18	WD Stanish	2013	Canada	RCT	41	39	Microfracture with BST-CarGel	Microfracture	35.1	37.2	18	14	NA	NA	12
19	E Basad	2010	Germany	RCT	40	20	MACI	Microfracture	33.0	37.5	15	3	7.0	7.0	24

20	E Solheim	2017	Norway	RCT	20	20	Microfracture	Mosaicplasty	35.0	31.0	6	6	4.0	4.0	180
21	S Bisicchia	2019	Italy	RCT	20	20	Microfracture with SVF	Microfracture	49.8	46.1	8	7	3.2	3.1	12
22	DBF Saris	2014	Netherlands	RCT	72	72	MACI	Microfracture	34.8	32.9	27	24	4.9	4.7	24
23	DBF Saris	2008	Netherlands	RCT	57	61	ACI-P	Microfracture	33.9	33.9	22	20	2.6	2.4	12
24	DBF Saris	2009	Netherlands	RCT	57	61	ACI-P	Microfracture	33.9	33.9	22	20	2.6	2.4	36
25	Z Qiao	2020	China	RCT	10	10	Microfracture	Microfracture with HA	62.3	59.7	7	5	4.0	4.0	12
						10		Microfracture with MSC		62.0		7		4.0	12
26	PD Nguyen	2016	Vietnam	RCT	15	15	Microfracture with SVF	Microfracture	58.6	58.2	12	12	NA	NA	18
27	H Lim	2021	Republic of Korea	RCT	43	46	Microfracture with MSC	Microfracture	55.3	54.4	28	30	4.9	4.0	60
28	M Venosa	2022	Italy	RCT	19	19	Microfracture with PRP	Microfracture with MSC	56.4	55.8	7	10	1.0	1.0	12
29	MS Shive	2014	Canada	RCT	34	26	Microfracture with BST-CatGel	Microfracture	34.3	40.1	12	12	2.4	2.0	60
30	Y Koh	2015	Republic of Korea	RCT	40	40	Microfracture with MSC	Microfracture	39.1	38.4	24	26	4.8	4.6	24
31	G Knutsen	2004	Norway	RCT	40	40	ACI-P	Microfracture	33.0	31.1	16	16	5.1	4.5	24
32	MS Kim	2017	South Korea	RCT	14	14	Microfracture	Microfracture with Collagen	55.7	55.4	0	1	2.9	3.6	12
33	MS Kim	2019	South Korea	RCT	48	52	Microfracture	Microfracture with Collagen	51.7	48.9	9	12	4.6	3.9	24
34	MS Kane	2018	United States	RCT	21	9	Neocart	Microfracture	41.4	38.8	2	3	2.2	1.7	60
35	C Ibarra	2021	United	RCT	24	24	MACI	Microfracture	33.7	35.8	7	10	1.9	1.7	72

36	Y	2019	Japan	RCT	7	4	Microfracture with MSC	46.3	4	0	3.0	4.4	12
	Hashimoto												
37	R Gudas	2006	Lithuania	RCT	28	29	AOT	24.3	10	12	2.8	2.7	36
38	R Gudas	2012	Lithuania	RCT	28	29	AOT	24.3	10	12	2.7	2.8	120
39	R Gudas	2005	Lithuania	RCT	29	28	Microfracture	24.3	12	10	2.8	2.7	36
40	J	2020	Germany	RCT	12	12	Microfracture	47.9	3	6	1.7	1.7	12
	Glasbrenner												
41	U Dasar	2016	Turkey	RCT	20	20	Microfracture	38.5	15	15	3.5	4.0	24
							Carbon fibre rod						
42	DC	2012	United States	RCT	21	9	NeoCart	39.0	2	3	2.8	2.5	24
	Crawford												
43	BJ Cole	2011	United States	RCT	9	20	Microfracture	32.7	4	6	3.4	2.7	24
							MACI						
44	JY Chung	2013	South Korea	RCT	24	12	Microfracture with BMAC	44.3	10	10	1.3	1.5	24

ACI: Autologous chondrocyte implantation; ACI-C: ACI with collagen cover; ACI-P: ACI with periosteal cover; AMIC: Autologous matrix induced chondrogenesis; BMAC: Bone marrow aspiration concentrate; CCP: Cultured chondrocyte pellet; HA: Hyaluronic acid; MACI: Matrix-induced autologous chondrocyte implantation; MEx: Microfracture; MSC: Mesenchymal stromal cell; NA: Not available; OAT: Osteochondral autograft/allograft transfer; PRP: Platelet-rich plasma; RCT: Randomized controlled trial; SVF: Stromal vascular fraction.

Table 2 Network meta-analysis summary and ranking of interventions based on the SUCRA scores

Follow-up	Outcome	Intervention	Coeffecient	Standard error	SUCRA ranking
1 yr	VAS	MFX-II	0.139	0.296	MFX-III > MFX-II > MFX-I ¹
		MFX-III	0.023	0.457	
		MFX-II	-2.296	2.835	MFX-III > MFX-II > MFX-I ¹
	KOOS	MFX-III	-2.296	5.775	
		MFX-II	-17.008	11.160	MFX-I > MFX-III > MFX-II ³
	Lysholm score	MFX-III	-5.660	4.427	
		MFX-II	2.782	1.811	MFX-III > MFX-II > MFX-I ¹
	IKDC score	MFX-III	4.339	2.228	
		MFX-II	4.257	4.543	MFX-II > MFX-I ¹
	Cincinnati score	MFX-III	0.383	0.312	
		MFX-II	1.860	1.770	MFX-III > MFX-II > MFX-I ¹
	MRI filling	MFX-III			

2 yr	MOCART score	MFX-II	11.950	7.419	MFX-III > MFX-II > MFX-I ¹
	Adverse events	MFX-III	30.700	14.168	
		MFX-II	-0.529	0.373	MFX-III > MFX-II > MFX-I ¹
	Failure events	MFX-III	-0.138	0.546	
		MFX-II	-0.520	0.777	MFX-II > MFX-I ¹
	VAS				I ¹
		MFX-II	0.377	0.452	MFX-III > MFX-II > MFX-I ¹
	KOOS	MFX-III	0.690	0.795	
		MFX-II	1.899	2.971	MFX-II > MFX-I ¹
	Lysholm score				I ¹
		MFX-II	0.550	6.952	MFX-II > MFX-I > MFX-III ³
	IKDC score	MFX-III	-19.560	9.814	
		MFX-II	4.548	4.545	MFX-III > MFX-II > MFX-I ¹
	Cincinnati score	MFX-III	7.947	9.405	
		MFX-II	-6.227	3.775	MFX-II = MFX-

	MRI filling	MFX-II	0.840	0.468	I ² MFX-II > MFX-III > MFX-I ³
		MFX-III	0.418	0.508	
	MOCART	MFX-III	10.600	9.281	MFX-III > MFX-I ¹
5 yr	VAS	MFX-III	1.900	1.917	MFX-III > MFX-I ¹
	IKDC score	MFX-III	3.000	2.121	MFX-III > MFX-I ¹

¹Newer generations better than older generations.

²Newer generations equal to older generations.

³Newer generations worse than older generations.

VAS: Visual Analog Scale; mFX: Microfracture; KOOS: Knee Osteoarthritis Outcome Scale; IKDC: International Knee Documentation Committee; MRI: Magnetic resonance imaging; MOCART: Magnetic resonance observation of cartilage repair tissue.

Table 3 Risk of bias for all the pairwise comparisons for functional outcome from the network assessed with Cochrane’s Confidence in network meta-analysis approach

Comparison	Number of studies	Within-study bias	Reporting bias	Indirectness	Imprecision	Heterogeneity	Incoherence	Confidence rating	Reasons for downgrading
MFx-I: MFx-II	7	Some concerns	Some concerns	No concerns	Major concerns	Some concerns	No concerns	Moderate	Imprecision in results
MFx-I: MFx-III	1	Some concerns	Some concerns	No concerns	Major concerns	Some concerns	No concerns	Moderate	Imprecision in results
MFx-II: MFx-III	1	Some concerns	Some concerns	No concerns	Major concerns	Some concerns	No concerns	Moderate	Imprecision in results

MFx: Microfracture.

13%

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8 Kristofer J. Jones, Benjamin V. Kelley, Armin Arshi, David R. McAllister, Peter D. Fabricant. "Comparative Effectiveness of Cartilage Repair With Respect to the Minimal Clinically Important Difference", The American Journal of Sports Medicine, 2019

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