Management of Focal Chondral Lesion in the Knee Joint

Seung-Suk Seo, MD, PhD, Chang-Wan Kim, MD, PhD and Dae-Won Jung, MD

Department of Orthopaedics, Busan Paik Hospital, College of Medicine, Inje University, Busan, Korea

Articular cartilage does not contain vascular, nervous and lymphatic tissue and chondrocytes hardly participate in the healing or repair process of chondral tissue because of being surrounded by plenty of extracellular matrix. Therefore, the injury to articular cartilage frequently requires an operative treatment. The goal of surgical repair of articular cartilage is to regenerate nearly normal chondral tissue and prevent degenerative arthritis caused by the articular cartilage defect. Microfracture is a kind of cartilage repair procedure that makes a fibrin clot containing mesenchymal stem cells in the chondral lesion. Microfracture is a simple procedure but it has a disadvantage that the repaired tissue is fibrocartilage. Autologous chondrocyte implantation has an advantage that it implants fully differentiated chondrocytes to the lesion, which theoretically produces hyaline cartilage. Its disadvantages are that it is a two stage and a costly procedure. Osteochondral autograft transplantation is a one stage procedure and repairs the lesion with hyaline cartilage. But its limitation is the lack of donor site availability. Surgeons who understand the theoretical background, indications, surgical methods, rehabilitation, complications, and clinical course of cartilage repair procedures can achieve the goal of preventing degenerative arthritis.

Key words: Knee joint, Articular cartilage, Focal chondral lesion, Management.

Introduction

Articular cartilage is devoid of vascular, nervous and lymphatic tissue and chondrocytes are unable to participate in the healing or repair process of damaged tissue because of extracellular matrix that surrounds the cartilage cells. About 200 years ago, a Scottish doctor, William Hunter, documented that chondral lesions had been considered difficult to treat and heal from the time of Hippocrates. In cases where chondral defects involve the subchondral bone causing bleeding, mesenchymal stem cells in the chondral lesion. Microfracture is a simple procedure but it has a disadvantage that the repaired tissue is fibrocartilage. Autologous chondrocyte implantation has an advantage that it implants fully differentiated chondrocytes to the lesion, which theoretically produces hyaline cartilage. Unfortunately, fibrocartilage is biomechanically inferior to hyaline cartilage and eventually results in osteoarthritis. Therefore, the goal of treatment for articular cartilage defects is to regenerate hyaline-like cartilage to prevent osteoarthritis. Total knee replacement can be helpful for patients with advanced age and severe osteoarthritis, but there are only a few options for young osteoarthritic patients. For successful focal chondral defect treatment, the prevalence of articular cartilage damage, structures and characteristics of the articular cartilage, post-injury responses, and scientific bases and clinical outcomes of various treatment methods should be well understood.

Treatments

The goal of focal chondral defect treatment is to enable patients to return to normal activities or active lifestyle through pain relief and joint function improvement. The treatment decision should be based on the patient's activity level, age, cause, size and depth
of defects, and presence of combined defects. Available treatment options include conservative treatment, surgical treatment for symptom relief or articular cartilage restoration.

1. Conservative Treatment
Conservative treatment for chondral defects of the knee can be effective for pain relief, but it cannot be used for articular cartilage restoration. Craig et al. suggested that conservative treatment can be an option when mild pain is present or the risk of surgery is greater than its benefit. Messner and Maletius followed up 28 patients with isolated chondral damage of the knee for 14 years and concluded that conservative treatment was not helpful in preventing the progress of the damage: the patients had excellent or good clinical results, but radiographic examination revealed abnormal findings in ≥50% of the patients. Conservative treatment includes the use of non-steroid anti-inflammatory drugs (NSAIDs), pain killers, hormones (estrogen and growth hormone), chondroprotective agents (glucosamine & chondroitin phosphate and omega-3), intraarticular injections of steroids or hyaluronic acid, weight loss to reduce the load on the knee joint, braces, and physical treatment. Unfortunately, these methods can be useful for symptom relief only not for restoring structural integrity of the articular cartilage.

2. Surgical Treatment
The purpose of surgical treatment is to improve symptoms and prevent degenerative changes by achieving structural and biomechanical restoration of the articular cartilage. Surgical treatment methods can be broadly divided into arthroscopic lavage and debridement, cell-based therapy (subchondral bone stimulation for chondral tissue differentiation or culture and implantation of chondrocytes), and tissue-based therapy (osteochondral autograft transplantation or osteochondral allograft transplantation). The advantages/disadvantages of each method, size, location, and depth of a lesion, and the patient’s age and activity level should be assessed to determine an appropriate treatment method. However, the two most important factors that should be considered are the cause and characteristics of chondral defects. Chondral lesions can be either focal or degenerative. For the treatment of focal lesions, sufficient debridement should be performed to maintain the adjacent area in the articular cartilage healthy for successful structural and biomechanical restoration. In contrast, for degenerative lesions where the defective and transitional area is wide, a sufficient debridement may restrict subsequent treatment options or cause unfavorable results. In addition, poor cell/tissue regeneration ability may result in less than satisfactory outcome after surgery. Therefore, the cause and characteristics of chondral lesions should be taken into consideration in performing surgical treatment.

1) Arthroscopic lavage and debridement
There has been a transition from open to arthroscopic lavage and debridement of chondral lesions. Arthroscopic lavage is to remove inflammatory mediators that may be responsible for joint effusion and loose cartilage and collagen debris. Debridement of articular cartilage (chondroplasty) is a procedure for removing unstable cartilage fragments or margins of the cartilage that may cause joint impingement with a curette or a shaver in order to alleviate joint pain and prevent additional articular cartilage destruction. Jackson et al. observed clinical improvements in 80% of their patients at 3.5 years after arthroscopic debridement and correlated degenerative changes with clinical outcome. However, Kirkley et al. reported that arthroscopic surgery for knee cartilage defects provided no additional benefit to optimized physical or medical therapy. Arthroscopic repair can be helpful for preventing the progress of delamination of articular cartilage, but there is controversy over its influence on the long-term longevity of the articular cartilage.

2) Cell-based therapy
Cell-based therapy is a promising approach using the patient’s own cells for the treatment of chondral defects. There are marrow stimulating procedures and autologous chondrocyte implantation (ACI). Marrow stimulating procedures include abrasion arthroplasty, drilling, and microfracture. However, the former two are currently rarely performed because they have been associated with poor clinical outcome.

(1) Microfracture
Microfracture is an articular cartilage repair technique in which tiny fractures are made 2-4 mm apart from each other to cause bleeding in the subchondral bone and fibrin clots in the perforations release mesenchymal stem cells that would differentiate into chondrocytes.

Multi-potential mesenchymal stem cells can differentiate into fibrocartilage cells and chondrocytes and induce fibrocartilage or hyaline-like cartilage formation. Fibrocartilage contains more collagen and less proteoglycans compared to hyaline cartilage. It is composed of more type I collagen than type II collagen. Type I collagen has lower compressive strength, elasticity, and wear resistance compared to type II collagen. Accordingly, fibrocartilaginous repair eventually results in failure
under repeated mechanical stress. In addition, the number of mesenchymal stem cells is small and tends to decrease over time. In spite of these disadvantages, microfracture has become a preferred procedure because it does not cause damage to other normal regions and is easy to perform and relatively economical.

Microfracture can be used for unstable or full-thickness (Outerbridge grade 3 or 4) focal chondral defects or degenerative arthritis with good knee alignment. The procedure is contraindicated for patients with inflammatory arthritis, lower limb malalignment, partial-thickness (Outerbridge grade 1 or 2) chondral defects, or reluctance to participate in rehabilitation. There are other considerations that should be taken into account. The older the patient is, the poorer the treatment outcome. Traumatic lesions have been associated with better treatment results compared to degenerative lesions. Theoretically, microfracture can be performed regardless of the size of a lesion. However, Steadman et al. described that lesions larger than 400 mm² responded worse to the treatment. In addition, the higher the body mass index, the poorer the treatment outcome. The procedure consists of debridement and drilling. Debridement should be performed thoroughly. All loose cartilage should be completely removed to make the lesion surrounded by healthy cartilage and form perpendicular edges to create a well shouldered lesion (Fig. 1). This is intended to make marrow clots firmly adhere to the lesion and reduce direct load across the lesion for stable recovery. The remaining calcified cartilage layer should be completely removed because it inhibits attachment of repair tissue. During this procedure, care should be taken to avoid excessive damage to the subchondral bone. Subchondral bone drilling should be performed in a centripetal manner from the margin to the center of the lesion. Microfracture holes should be placed 3-4 mm apart from each other and 4-5 mm in depth (Fig. 1). An appropriate depth can be determined by observing fat droplets that is released from the marrow through the microfracture holes while controlling the perfusion pressure (Fig. 1).

Different postoperative rehabilitation regimens are adopted according to the location of a lesion. For femorotibial joint lesions, continuous passive motion exercise is initiated immediately after surgery. Steadman et al. reported that full passive range of motion could be obtained after 6 to 8 weeks of toe touch weight bearing and continuous passive motion exercise that had been started immediately after surgery. For femoropatellar joint lesions, care should be taken not to apply shear force on the lesions using a brace that allows for 0-20 degrees of joint movement. On the other hand, Marder et al. suggested that postoperative physical treatment or weight bearing does not

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**Fig. 1.** Surgical procedure of microfracture. (A) Unstable cartilage flap and calcified cartilage bed is debrided with open curette. (B) It is important to debride the calcified cartilage layer and make a well-contained pocket surrounded healthy cartilage (well-shouldered). (C) Subchondral bone is punctured with an awl. (D) Microfracture is circumferentially performed from periphery to center. (E) The penetration of subchondral bone is 3 to 4 mm deep and apart. (F) Arthroscopic photograph showing the final step of microfracture. (G) Mesenchymal blood egress from bone marrow through subchondral holes. (H) It is important for tissue regeneration to keep the mesenchymal clot in the defect.
have to be avoided when lesions are “well shouldered” during operation because the repair tissue is not affected by weight bearing based on their observation that aggressive physical treatment did not influence the treatment outcome.

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Peterson et al.\textsuperscript{22} observed good to excellent results in 50 out of 61 patients at two years after ACI and in 51 out of 60 patients at eleven years after ACI. A second-look arthroscopy showed a normal appearance of the cartilage in 90\% of the 8 patients and hyaline cartilage was confirmed by histological examination in most of the patients. Factors that could influence the outcome of ACI have also been addressed in many studies. Minas et al.\textsuperscript{23} reported that prior procedures using marrow stimulating techniques such as microfracture increase the failure rates of ACI. Mithoefer et al.\textsuperscript{24} observed poor clinical results in patients with large or multiple cartilage lesions. Saris et al.\textsuperscript{25} documented that the longer the duration of symptoms before ACI was, the poorer the clinical results. Knutsen et al.\textsuperscript{26} noted that patients under 30 years of age with high activity had good clinical results after ACI. These studies indicate that various factors should be taken into consideration to achieve satisfactory outcome after the procedure.

The first generation ACI has often been associated with periosteal complications. Kreuz et al.\textsuperscript{27} reported that clinical improvement was observed in most of the patients after ACI, but MRIs revealed periosteal hypertrophy in 28\% of the total patients and in 50\% of the patients with ACI for chondral defects in the knee. Seo et al.\textsuperscript{28} reported that ACI for chondral defects of the femoral condyle resulted in clinical improvement, but the International Cartilage Repair Society (ICRS) assessment revealed abnormal findings in 60\% and graft hypertrophy in 21\% of the patients (Figs. 3, 4).

The first generation ACI has been replaced by the second and then the third generation ACI to prevent periosteal complications and obtain structurally and biomechanically superior tissue. The second generation ACI is a procedure in which a synthetic collagen membrane instead of a periosteal flap is sutured to the defect and chondrocytes are injected. The third generation ACI is a procedure in which chondrocytes are cultured in a biodegradable scaffold that will be implanted in the defect. Although various studies have addressed the results

Fig. 2. Surgical procedure of the 1st generation autologous chondrocyte implantation. (A) Outerbridge 4 lesion in the medial femoral condyle. (B) Debridement of the calcified cartilage layer and unstable chondral flap. (C) Defect size is measured with a sterile paper. A 2 mm oversized template is needed. (D) Periosteal flap is excised from the proximal medial tibia. (E) The periosteal flap watertight covers the defect. (F) Chondrocyte suspension is injected to the defect through a plastic 18-gauge angiocath needle.
of each procedure, more studies should follow to confirm those results\(^{29-35}\).

ACI is useful for maintaining good clinical outcome for a long-term period in cases of focal cartilage defects, but the disadvantages include that it requires two separate procedures for chondrocyte collection and implantation and joint excision is unavoidable in most cases during implantation. With the advent of the second and third generation ACI technique, the procedure can be performed arthroscopically and has become easier to perform. We expect further improvements will be made in the future.

3) Tissue-based therapy

Tissue-based therapy includes osteochondral autograft transplantation, osteochondral allograft transplantation, and tissue engineered scaffold implantation. The advantages of these procedures are treating defects in one stage, promoting rapid return to daily living activities and sports with use of biomechanically healthy tissue, and maintaining postoperative results for a long-term period due to hyaline cartilage repair. Since tissue engineered scaffold implantation can be used in limited clinical settings and accordingly only a small number of clinical results have been reported, we describe the former two methods in this review article.
(1) Osteochondral autograft transplantation (OAT, mosaicplasty)

OAT can be a solution for hyaline repair of chondral defects. The advantages of this technique include: 1) it is a one-stage procedure unlike ACI; 2) it can be performed arthroscopically for small lesions; 3) osteochondral plugs can be obtained with ease; 4) it is a tissue-based therapy that allow earlier rehabilitation compared to cell-based therapy; 5) it can be performed at a lower cost compared to ACI; 6) the lesion is covered by hyaline cartilage; and 7) it results in few complications. One of the disadvantages of the procedure is that it cannot be applied for large lesions due to limited donor site availability. For large lesions, multiple osteochondral plugs are necessary, but it is difficult to coordinate the height and direction of the implanted osteochondral plugs with the adjacent native cartilage. The gaps between osteochondral plugs and between the plugs and the surrounding cartilage are filled with fibrocartilage. In addition, posttraumatic arthritis may develop in the patellofemoral joint due to graft harvest.

The primary indication for OAT is a symptomatic anterior cartilage defect (Outerbridge stage 3 or 4) that is 1.0-4.0 cm² in size in patients less than 45 years of age. The contraindications include patients above 50 years of age, larger than 8.0 cm² lesions, moderate or severe osteoarthritis, inflammatory arthritis, lack of appropriate donor site, and noncompliance with rehabilitation. Besides, lower limb alignment and combined ligament injuries should also be assessed and treated prior to OAT.

The procedure consists of defect preparation, graft harvesting with a tubular chisel and graft removal from the chisel, and transplantation of the graft. On defect preparation, unstable cartilage is removed and the size of a lesion is measured to determine the size, number, and arrangement of osteochondral plugs. The most common donor sites are the superolateral aspect of the femoral intercondylar notch and the medial/lateral margin of the femoral trochlea. The osteochondral plug can be harvested using a tubular chisel and the appropriate length is 15 mm. It is important to direct the tubular chisel perpendicular to the articular surface to obtain a graft that has a level articular surface. The graft is ejected from the chisel by tapping the osseous tissue, not the articular surface. Next is the transplantation of the osteochondral plug into the defect, which consists of drilling, dilation, and delivery. An appropriately sized drill guide is placed perpendicular to the walls of the lesion and the same diameter drill bit is introduced to create a tunnel in the lesion. The tunnel

Fig. 5. Surgical procedure of osteochondral autograft transplantation. (A) Arthroscopic determination of the number and size of grafts needed after debridement of cartilage lesion. (B) Open procedure. (C) Harvesting the osteochondral plug with a tubular chisel from the lateral supracondylar ridge. The tubular chisel must be perpendicularly located to the chondral surface. (D) Introduction of the osteochondral plug through a drill guide. (E) Harvested osteochondral plugs. Size and length of plugs are marked on the wet gauze. (F) Focal cartilage defect reconstructed with multiple osteochondral plugs. It is important to make a congruent surface with the adjacent cartilage.
should be approximately 2 mm longer than the graft length and widened with a dilator of the same diameter. The graft is gently tapped into the hole using a delivery tamp to avoid chondrocyte death. When several grafts need to be implanted, the procedure is repeated for each graft to increase mechanical stability. It is crucial to ensure that the articular surface of the graft is level with the adjacent articular surface (Fig. 5). In a biomechanical study conducted by Koh et al., when a graft was placed slightly deeper than the adjacent articular cartilage, the contact pressure was normal, whereas 2 mm elevation of a graft resulted in an approximately 50% increase in the pressure. In an animal study, cartilage necrosis and fibrous overgrowth were observed when grafts were placed 2 mm deeper than the adjacent articular cartilage.

OAT allows rapid rehabilitation. Continuous passive motion and straight leg raising are performed immediately after surgery. Weight bearing is allowed depending upon the number of the implanted grafts. Generally, non-weight bearing or tip-toe walking with use of a crutch is permitted for 2 postoperative weeks. Partial weight bearing is allowed for two or four postoperative weeks and full weight bearing is possible from the fourth or sixth postoperative week.

The clinical outcome of OAT is relatively favorable. Many studies have shown that the technique was effective in 76-93% of the patients in achieving clinical improvement. However, there are some factors that could influence the clinical outcome. Jacob et al. noted high complication/reoperation rates in patients with large-sized lesions. Seo et al. reported that although OAT resulted in good clinical outcome, the improvement was relatively less remarkable in patients who were ≥30 years of age or had ≥4.0 cm² lesions (Fig. 6).

(2) Osteochondral allograft transplantation

Osteochondral allograft transplantation can be performed regardless of the size of a lesion and donor site availability and morbidity. The procedure can be employed for treating large lesions, ≥10 cm² in size. The drawbacks of the procedure include the difficulty of obtaining grafts in a timely manner,
high cost, and the possibility of immune rejection response and disease transmission. According to the graft type, there are shell type and deep type allografts. Shell type grafts include <1 cm subchondral bone and deep type grafts include the deeper layer of subchondral bone. According to the graft preservation method, there are fresh allografts, cryopreserved frozen allografts, and fresh frozen allografts. Fresh allografts are stored at 4°C and implanted within 1 week of harvesting. Compared to other grafts, fresh allografts have relatively high risk of immune rejection response or disease transmission because it is difficult to completely remove donors’ blood from the grafts. Cryopreservation is a process where allografts are preserved in glycerol or dimethyl sulfoxide to minimize chondrocyte death and maintain chondrocyte viability. It is effective for long-term tissue preservation and prevention of immune rejection response and disease transmission, but it has been associated with low survival rates of chondrocytes. Regarding the fresh frozen preservation, grafts are frozen at -80°C immediately after harvesting and accordingly carry low risk of infection or immune response. This procedure costs less than Cryopreservation does. However, chondrocytes are destroyed during freezing and biomechanical properties of extracellular matrix deteriorate over time.

The ideal indications for osteochondral allograft transplantation include traumatic osteoarthritis combined with extensive cartilage defects or subchondral bone defects in active patients, osteochondritis dissecans, and focal avascular necrosis. The contraindications include moderate or severe degenerative arthritis, inflammatory arthritis such as degenerative arthritis, and steroid-induced osteonecrosis.

Gross et al. reported long-term follow-up results of the use of fresh osteochondral allografts for post-traumatic knee defects: the survival rate was 95%, 80%, and 65% at 5, 10, and 15 years after surgery, respectively. According to Ghazavi et al. osteochondral allograft transplantation resulted in successful outcome in 85% and failure in 15% of the cases at 7.5 years after surgery. Regarding the factors that influence the osteochondral allograft transplantation, Ghazavi et al. attributed failure to bipolar tibial and femoral defects, malaligned knees, and workers’ compensation status. Osteochondral allograft transplantation can be advantageous in maintaining satisfying results for isolated extensive chondral or osteochondral defects that cannot be managed with OAT or ACI.

Conclusions

Chondral defects are difficult to treat. Various recent studies have introduced new repair techniques and reported clinical results. However, there is no consensus regarding which method is superior to the others. Factors that should be taken into consideration in determining a surgical method for focal chondral defects include the cause of the defect, concomitant defects, the patient's age and activity level, knee alignment, defect size, cost, and risk of surgery. Degenerative defects cannot be properly managed with surgical repair of articular cartilage. For patients with low activity, arthroscopic debridement is the only...
option for knee defects irrespective of the size. Concomitant defects on the affected knee or malalignment should be treated before surgical articular cartilage repair. For lesions that are 1-2 cm$^2$ in size, microfracture or arthroscopic debridement can be a treatment of choice in patients with low activity whereas OAT is effective in patients with high activity. ACI can be a solution when patients with high activity lack sufficient donor sites or microfracture or OAT resulted in failure. For lesions larger than 4 cm$^2$ in size, ACI is the most preferred method and bone grafting should be additionally performed when bone loss is present. Osteochondral defects larger than 10 cm$^2$, focal osteonecrosis, and post-traumatic osteochondral defects are treated with osteochondral allograft transplantation (Fig. 7).

Articular cartilage does not respond well to healing. Damage to the articular cartilage eventually degenerates into arthritis. The goal of current techniques for chondral defects is to prevent the degenerative process by regenerating hyaline like cartilage. Among various methods that have been introduced to achieve this goal, the optimal treatment choice should be based on the understanding of the theoretical backgrounds, indications, surgical technique, rehabilitation, complications, and clinical course of the treatment.

References

1. Buckwalter JA, Mankin HJ. Articular cartilage. Part II: degeneration and osteoarthritis, repair, regeneration, and transplantation. J Bone Joint Surg Am. 1997;79:612-32.
2. Shah MR, Kaplan KM, Meislin RJ, Bosco JA 3rd. Articular cartilage restoration of the knee. Bull NYU Hosp Jt Dis. 2007;65:51-60.
3. Curl WW, Krome J, Gordon ES, Rushing J, Smith BP, Poehling GG. Cartilage injuries: a review of 31,516 knee arthroscopies. Arthroscopy. 1997;13:456-60.
4. Craig W, David JW, Ming HZ. A current review on the biology and treatment of the articular cartilage defects (part I & part II). J Musculoskelet Res. 2003;7:157-81.
5. Messner K, Maletius W. The long-term prognosis for severe damage to weight-bearing cartilage in the knee: a 14-year clinical and radiographic follow-up in 28 young athletes. Acta Orthop Scand. 1996;67:165-8.
6. Guettler JH, Demetropoulos CK, Yang KH, Jurist KA. Osteochondral defects in the human knee: influence of defect size on cartilage rim stress and load redistribution to surrounding cartilage. Am J Sports Med. 2004;32:1451-8.
7. Jackson RW, Gilbert JE, Sharkey PF. Arthroscopic debride-
19. Mithoefer K, Williams RJ 3rd, Warren RF, Wickiewicz TL, Marx RG. High-impact athletics after knee articular cartilage repair: a prospective evaluation of the microfracture technique. Am J Sports Med. 2006;34:1413-8.

20. 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;331:889-95.

21. Jones DG, Peterson L. Autologous chondrocyte implantation. Instr Course Lect. 2007;56:429-45.

22. Peterson L, Minas T, Brittberg M, Nilsson A, Sjogren-Jansson E, Lindahl A. Two- to 9-year outcome after autologous chondrocyte transplantation of the knee. Clin Orthop Relat Res. 2000;(374):212-34.

23. Minas T, Gomoll AH, Rosenberger R, Royce RO, Bryant T. Increased failure rate of autologous chondrocyte implantation after previous treatment with marrow stimulation techniques. Am J Sports Med. 2009;37:902-8.

24. Mithofer K, Peterson L, Mandelbaum BR, Minas T. Articular cartilage repair in soccer players with autologous chondrocyte transplantation: functional outcome and return to competition. Am J Sports Med. 2005;33:1639-46.

25. 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 1:105-95.

26. Knutsen G, Engebretsen L, Ludvigsen TC, Drosgot JO, Grontvedt 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;86:455-64.

27. Knutsen G, Engebretsen L, Ludvigsen TC, Drosgot JO, Grontvedt 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;86:455-64.

28. Seo SS, Ha DJ, Kim CW, Kim HJ, Moon SW. Autologous chondrocyte implantation for treating chondral defects of the femoral condyle. J Korean Knee Soc. 2009;21:276-85.

29. Steinwachs M, Kreuz PC. Autologous chondrocyte implantation in chondral defects of the knee with a type I/III collagen membrane: a prospective study with a 3-year follow-up. Arthroscopy. 2007;23:381-7.

30. 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;13:203-10.

31. Tohyama H, Yasuda K, Minami A, Majima T, Iwasaki N, Muneta T, Sekiya I, Yagishita K, Takahashi S, Kurokouchi K, Uchio Y, Iwasa J, Deie M, Adachi N, Sugawara K, Ochi M. Atelocollagen-associated autologous chondrocyte implantation for the repair of chondral defects of the knee: a prospective multicenter clinical trial in Japan. J Orthop Sci. 2009;14:579-88.

32. Nehrer S, Dorotka R, Domayer S, Stelzeneder D, Kotz R. Treatment of full-thickness chondral defects with hyalograph C in the knee: a prospective clinical case series with 2 to 7 years' follow-up. Am J Sports Med. 2009;37 Suppl 1:81S-7S.

33. Kim MK, Choi SW, Kim SR, Oh IS, Won MH. Autologous chondrocyte implantation in the knee using fibrin. Knee Surg Sports Traumatol Arthrosc. 2010;18:528-34.

34. 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;87:640-5.

35. 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;38:924-33.

36. Hangody L, Rathonyi GK, Duska Z, Vasarhelyi G, Fules P, Modis L. Autologous osteochondral mosaicplasty. Surgical technique. J Bone Joint Surg Am. 2004;86 Suppl 1:65-72.

37. Duchow J, Hess T, Kohn D. Primary stability of press-fit-implanted osteochondral grafts. Influence of graft size, repeated insertion, and harvesting technique. Am J Sports Med. 2000;28:24-7.

38. Whiteside RA, Jakob RP, Wyss UP, Mainil-Varlet P. Impact loading of articular cartilage during transplantation of osteochondral autograft. J Bone Joint Surg Br. 2005;87:1285-91.

39. Koh JL, Wirsing K, Lautenschlager E, Zhang LO. The effect of graft height mismatch on contact pressure following osteochondral grafting: a biomechanical study. Am J Sports Med. 2004;32:317-20.
40. Huang FS, Simonian PT, Norman AG, Clark JM. Effects of small incongruities in a sheep model of osteochondral autografting. Am J Sports Med. 2004;32:1842-8.
41. Jakob RP, Franz T, Gautier E, Mainil-Varlet P. Autologous osteochondral grafting in the knee: indication, results, and reflections. Clin Orthop Relat Res. 2002;(401):170-84.
42. Seo SS, Kim CW, Ha DJ, Choi JS, Kim HJ, Lee CR. Autogenous osteochondral grafting for treating osteochondral defect of the femoral condyle of the knee joint. J Korean Orthop Assoc. 2009;44:301-10.
43. Chu CR, Convery FR, Akeson WH, Meyers M, Amiel D. Articular cartilage transplantation. Clinical results in the knee. Clin Orthop Relat Res. 1999;(360):159-68.
44. Bugbee WD, Convery FR. Osteochondral allograft transplantation. Clin Sports Med. 1999;18:67-75.
45. Enneking WF, Mindell ER. Observations on massive retrieved human allografts. J Bone Joint Surg Am. 1991;73:1123-42.
46. Enneking WF, Campanacci DA. Retrieved human allografts: a clinicopathological study. J Bone Joint Surg Am. 2001;83:971-86.
47. Hennig A, Abate J. Osteochondral allografts in the treatment of articular cartilage injuries of the knee. Sports Med Arthrosc. 2007;15:126-32.
48. Gross AE, Shasha N, Aubin P. Long-term followup of the use of fresh osteochondral allografts for posttraumatic knee defects. Clin Orthop Relat Res. 2005;(435):79-87.
49. Ghazavi MT, Pritzker KP, Davis AM, Gross AE. Fresh osteochondral allografts for post-traumatic osteochondral defects of the knee. J Bone Joint Surg Br. 1997;79:1008-13.