The menisci and articular cartilage of the knee have a close embryological, anatomical and functional relationship, which explains why often a pathology of one also affects the other.

Traumatic meniscus tears should be repaired, when possible, to protect the articular cartilage.

Traumatic articular cartilage lesions can be treated with success using biological treatment options such as microfracture or microdrilling, autologous chondrocyte transplantation (ACT), or osteochondral transplantation (OCT) depending on the depth and area of the lesion.

Degenerative cartilage and meniscus lesions often occur together, and osteoarthritis is already present or impending. Most degenerative meniscus lesions should be treated first conservatively and, after failed conservative treatment, should undergo arthroscopic partial meniscus resection. Degenerative cartilage lesions should also be treated conservatively initially and then surgically; thereby treating the cartilage defect itself and also maintaining the axis of the leg if necessary.

Tears of the meniscus roots are devastating injuries to the knee and should be repaired e.g. by transtibial re-fixation.

The clinical role of ‘ramp’ lesions of the meniscus is still under investigation.

**Keywords:** arthroscopic partial meniscectomy; cartilage; conservative treatment; meniscus; meniscus ‘ramp’ lesions; meniscus repair; meniscus roots; osteoarthritis

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### Anatomy, function and epidemiology

The close anatomical and functional relationship of menisci and articular cartilage, in many parts very similar organs, starts during their embryological development. Both articular cartilage and menisci, medial and lateral, originate from the interzone cells. Menisci, also known as semilunar cartilages, cover a large part of the cartilage of the tibial plateau (Fig. 1), whereby the cartilage of the femoral condyle slides during movement of the knee above the menisci and the uncovered chondral parts of the tibial plateau. Menisci have several fixation points. The roots of the meniscus are the main stabilizers. Each human meniscus has an anterior and a posterior root. The roots are ligamentous structures connecting the meniscus horns to the tibial plateau. In addition, the posterior root of the lateral meniscus has two inconsistently existing stabilizers: the anterior and posterior meniscofemoral ligaments (Humphrey and Wrisberg), originally described by the German anatomist Josias Weitbrecht in 1742 (Fig. 2). However, there are more structures that provide stability to the menisci, for example, the transverse ligament connecting both anterior meniscus horns and the coronary ligament including the ‘meniscus ramp’. The meniscus ramp is the part of the coronary ligament which connects the posterior horn of the medial meniscus to the tibial head.

Histologically, one of the main components of both cartilage and menisci is collagen: menisci contain mainly type I and articular cartilage type II collagen. Both are also dependent on collagen type III, which modulates the thickening of the aberrant fibres of type I collagen in menisci and type II collagen in cartilage.

Besides their anatomical relationship, cartilage and menisci also have a close functional relationship. Menisci create congruity in the knee joint, allowing the cartilage of the femoral condyles to slide smoothly over the menisci and the tibial cartilage. Cartilage primarily takes compressive loads. Menisci also take compressive loads but mainly in Cooper zone 3. Furthermore, they support proprioception of the knee joint, providing stability, and lubrication. However, the main function of menisci is to convert
Tibiofemoral load into hoop stress through its longitudinal fibres, connecting the anterior and posterior meniscus roots. Disturbance in the structure of the menisci affects the distribution of forces and pressure load on the articular cartilage, leading to an early onset of osteoarthritis (OA). This close relationship between menisci and articular cartilage has also been demonstrated through the impact of degenerative and trauma-related meniscal injuries on the articular cartilage. Degenerative meniscus lesions, mostly horizontally orientated, develop over time. Patients with these kinds of lesions have not experienced a trauma which is sufficient to tear their meniscus. These lesions can extend into flaps or complex lesions and are often associated with meniscus extrusion and osteoarthritis. The prevalence of degenerative meniscus lesions increases with age. Among 50 to 59-year-old females, 19% of patients had a degenerative meniscus lesion on magnetic resonance imaging (MRI). Among 70 to 90-year-old male patients, 56% had a degenerative meniscus lesion. About two-thirds of these lesions affect the medial meniscus. The prevalence of degenerative meniscus lesions increases with higher degree of OA. In a group of patients with OA of Kellgren and Lawrence (K&L) III or IV, 95% of patients had a damaged meniscus. Patients with meniscus extrusion are even more prone to develop early OA (medial tibial plateau: (a) 2.1% per annum with extrusion vs. (b) 1.5% without extrusion; lateral tibial plateau: (a) 2.6% vs. (b) 1.6%, respectively). This extrusion can occur in patients with degenerative meniscus lesions and with meniscus root tears. Besides degenerative meniscus lesions, there are also traumatic meniscus tears. These tears are mainly vertically orientated, for example, longitudinal (including bucket-handle) and radial tears. In general, patients report a significant trauma and are younger compared with patients with degenerative tears. Additionally, traumatic tears often occur concomitantly with ligament tears of the ipsilateral knee, e.g. a rupture of the anterior cruciate ligament (ACL). About 6% of injured knees sustain a traumatic meniscus tear resulting in 0.5 to 0.7 acutely injured menisci per 1000 individuals per year. The numbers of affected medial and lateral menisci in patients with a concomitant ACL tear vary significantly between studies. In general, it is accepted that in acutely injured knees with an ACL tear, lateral meniscus tears occur more frequently. By contrast, in chronically ACL insufficient knees, medial meniscus tears are more common. A similar discrimination is seen with meniscus root tears. Traumatic root tears mainly affect the posterolateral root in young patients with ACL tears; and, on the other hand, the degenerative meniscus root lesions including insufficiency (Fig. 3) mainly affect the posteromedial root in...
older patients (often aged > 50 years), without a significant trauma but often associated with degenerative cartilage lesions of the medial tibiofemoral compartment (Fig. 3).

Similarly, articular cartilage defects are different in traumatic and degenerative lesions. To classify them, the classification of the International Cartilage Regeneration and Joint Preservation Society (ICRS) has been the widely accepted. To classify osteoarthritis of the knee, a rather general joint failure not just cartilage degeneration, the Kellgren and Lawrence and the Ahlbäck classifications have been widely accepted.20–22 Osteoarthritis is defined as a set of conditions that affect diarthroses, characterized through stress manifested at a cellular level and degradation of the extracellular matrix (ECM), promoted by injuries at both micro and macro level which activate a pathological repair response from the immune system, including the pro-inflammatory pathway.23 Another definition proposed by the American College of Rheumatic and Therapeutic Criteria states that OA is a group of conditions that are heterogenic, which advances to joint-related symptoms and signs caused by the damaging of the articular cartilage structure alongside the damaging of the subsequently subchondral bone. What must also be understood is that illness is not the same as disease. Disease is characterized by signs and objectively quantified modifications, which in OA cases are: deformity, crepitus, restriction of movement and osteophytes. Illness refers to the subjective symptoms that a patient feels, such as mechanically characterized pain, joint stiffness and functional impairment. There are speculations that the threshold for clinical manifestation of OA is around 20% of joint affection.23 The pathological abnormalities progress from a molecular stage through an anatomical one and end with the physiological alterations. Molecular abnormalities such as altered proteins, metabolites and changes in the genomics can be demonstrated in early OA, long before any relevant modifications can be seen by X-rays, ultrasonography, MRI, and Positron emission tomography (PET) scans, during the so-called ‘silent period’ of time in which the patient also does not display any symptoms. Possible relevant markers that have been identified so far include: free type II collagen, urinary C-telopeptide of collagen II, racemized protein epitopes and deaminated protein epitopes. Also, the biochemical structure of the OA-affected articular cartilage differs from a normal one, creating a possible field for future research in this area. Anatomical abnormalities can be ascertained using different diagnostic modalities such as radiography, ultrasonography, MRI and PET scans. Through them the fibrillation of cartilage, loss of proteoglycans, apoptosis or proliferation of chondrocytes, malalignment of the joint, narrowing of joint space, osteophyte formation, presence of bone cysts and subchondral sclerosis can be observed. In the case of early OA, MRI has proven to be a great investigational tool. Thickening of cartilage, elevated T1rho signal, meniscal degeneration and bone shape modifications are some of the objective findings. Ultrasonography can be also useful in diagnosing early OA through findings such as synovitis and angiogenesis, which occur secondary to the release of cartilage degradation products into the joint. The very last changes that appear in the development of OA are of physiological nature and refer to the abnormal biomechanical forces that are exerted at the affected joint level and their effects on it.23 Cartilage lesions occur frequently. Out of 25,124 knee arthroscopies performed for different knee pathologies including meniscal tears, ACL reconstruction, acute trauma injuries, knee pain without known cause, etc.; 60% revealed different types of chondral lesions. From the aforementioned

Fig. 3 (A) Severe damage of the entire medial femoral condyle and tibial plateau with cartilage degradation. (B) The reason for this degradation is the stretched, insufficient posteromedial meniscus root. The root was stretched by an osteophyte lying underneath the root.
60%, 67% of the patients had focal chondral/osteochondral lesions, 29% displayed OA, 2% were diagnosed with osteochondritis dissecans and 1% had other lesions. The true incidence of chondral lesions though remains unknown due to the lack of symptoms. In terms of concomitant lesions, rupture of the medial meniscus ranks as first with 42%, followed by ACL injury with 36%. In terms of localization, the patellar surface and medial femoral condyle are the most common, at 36% and 34%, respectively.

Due to the fact that articular cartilage and menisci are subjected to the same forces and originate from the same embryonic interzone; a degenerative meniscus lesion tends to be associated with a pathological articular cartilage structure. There are findings that support the idea that structural changes inside the knee joint can start either from the meniscus or the cartilage. Degenerative lesions of the meniscus can progress to structural lesions of the subchondral bone such as: increase of the subchondral bone density, subchondral bone cysts and bone marrow lesions, alongside cartilage damage, due to a rise in peak stress. Subsequently, radiographic findings support the idea that lesions of the articular cartilage have a key role in developing lesions of the subchondral bone, like subchondral oedema. About 80% of middle-aged patients with meniscus lesions presented with typical symptoms such as general knee pain, pain during knee twisting and an untrustworthy knee (gives way). These symptoms matched the symptoms of patients with early radiological OA. As already mentioned above, degenerative meniscus lesions are often associated with articular cartilage wear. In a recent study, 43% of patients who had undergone meniscus-related arthroscopies presented with early or advanced knee OA. About 75% of patients had presented with large cartilage defects: 44% retropatellar, 34% on the medial femoral condyle, and 18% on the medial tibial plateau. Based on these findings and similar findings of others, it was postulated that the damage inflicted on the articular cartilage tends to be localized and not uniformly distributed across the femoral and tibial articular surfaces. More specifically, tears in the medial meniscus body had a damaging effect on the articular cartilage lining the tibial central and anterior subregion. The same effect was observed in cases of posterior horn lesion, but in the posterior tibial subregion. Out of all types of meniscus lesions, radial tears at the medial body were highly associated with loss of articular cartilage leading to OA. Interestingly, clinical syndromes are not mandatory in patients with degenerative meniscus lesions as data from the Framingham study showed, where prevalence of meniscus lesions was 31% in patients aged 50 to 90 years, and only 29% of these patients had knee complaints. These numbers increased significantly with increasing osteoarthritic changes. In patients with severe OA, up to 95% of patients had meniscus lesions (at least grade III) compared with 25% of patients without radiological signs of OA. In addition, there is a yet unknown link between tears in degenerated meniscus and hand OA, with a 25% prevalence of damaged meniscus in the knew of subjects with zero finger joints affected by OA, 32% prevalence in those with one to two finger joints with OA, and 47% in patients with three or more finger joints affected. This is surely an interesting finding which may provide clues for future studies regarding tears in degenerated menisci and generalized OA. Individual risk factors are, for example: female gender, older age, genetic pre-conditions, thickened intimal media of the carotid artery, tibial plateau fracture, ACL insufficiency and obesity. On the other hand, MRI examinations showed that weight loss...
resulting in a decreased BMI was associated with a slowdown of cartilage degeneration.36

**Treatment of meniscus and articular cartilage disorders**

Treatment recommendations differ depending on the pathologies. For degenerative meniscus lesions, a paradigm shift has happened over the last 10 years, suggesting primarily conservative treatment, and arthroscopy with partial meniscus resection not until a conservative course of treatment has failed. This is true for patients with and without knee osteoarthritis.37–41 However, patients should be informed that with increasing degenerative changes of the knee the success rate decreases. It is important to check the limb alignment and treat it if appropriate. A Western Ontario and McMaster Universities Osteoarthritis index (WOMAC) of > 40 points was identified as a predictor for failing conservative treatment and early arthroscopic partial meniscectomy.42 A recent European consensus agreed that the time period for conservative treatment should be at least three months.9 Thus far, no agreement exists about which kind of conservative treatment should be applied. Most studies used physiotherapy regimes or intra-articular corticosteroid injection. Perhaps a watch-and-wait approach is also an option. The role of non-steroidal anti-inflammatory drugs (NSAIDs) and others is unclear here, because the intake of analgesics was neither controlled nor monitored in these aforementioned randomized controlled trials (RCTs). These RCTs showed that conservative treatment is not inferior to partial meniscectomy; however, up to 35% of patients changed from conservative to surgical treatment during the trials (cross over). Some patients develop pain after arthroscopic partial meniscectomy, so that partial meniscus replacement might be an option, e.g. Actifit or collagen meniscus implant (CMI).43–45 These implants look initially similar to a meniscus and are tailored during surgery to fit into the meniscus defect. For both, an intact peripheral rim of the meniscus is critical to provide stability (Fig. 4).45 K&L changes should not exceed II° at the time of implantation. Clinical results have been promising, including apparently stopping or at least slowing down progression of OA.46 However, these implants have never been generally accepted because of considerable reported problems. Reasons for failure were a considerable resorption rate or decrease in size (Fig. 5), long rehabilitation and pending issues with availability because of regulatory authorities e.g. United States Food and Drug Administration (US FDA) or financial issues of the companies.46–48

If the peripheral rim is interrupted, a complete meniscus replacement is necessary. Traditionally, a human meniscus allograft has been used in this circumstance. However, because of its high price, legal and administrative issues in many countries, rather small improvement in patient-reported outcome measures (PROMs), return to just low-impact activities in general as well as its questionable prevention of a further progress of OA, alternative solutions have been explored.49–52 One of these recent solutions is NuSurface, a free-floating, ring-shaped, polycarbonate-urethane meniscus implant (Fig. 6).47,53,54

By contrast, the primary choice of treatment for traumatic meniscus tears is arthroscopic repair.55 Self-induced repair rarely happens, except for small longitudinal tears of the posterior horn of the lateral meniscus in patients with a concomitant ACL repair or reconstruction. A biomechanical study showed that a longitudinal tear in the medial meniscus body decreased the in situ meniscus force under an axial load, supporting the necessity of meniscus repair to avoid articular cartilage damage.56 For
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For a long time, the treatment of choice for traumatic meniscus tears was complete or partial resection, until recent studies showed the superiority of repair vs. partial meniscectomy with regard to the development of OA and return to the pre-injury level of activity. Tears should be repaired even if they are long or chronic. In cases where an ACL reconstruction is not concomitantly performed, the medullary cavity should be opened, e.g. three holes into the femur anteriorly to the ACL attachment utilizing a microfracture awl or a drill bit. General opinion is that longitudinal tears of Cooper zones two to zero should be repaired. This is true for men and women achieving similar results after five years of meniscus repair. Repair is also the treatment of choice in patients with meniscus root tears; both degenerative and traumatic. The most commonly used technique is a transtibial root repair, and a biomechanical study suggested a two-tunnel technique (Fig. 7). The sutures should be placed into the meniscus horns. In cases with a concomitant ACL reconstruction, on the tibia the sutures of the meniscus root should not be pulled through the tibial ACL tunnel but through a separate tunnel in order to achieve physiological tension of the repaired meniscus (Fig. 7).

For degenerative meniscus root tears, it is important to pay attention to the grade of OA and leg alignment. Repair of roots is superior to conservative treatment and partial meniscectomy in knees with K&L grades I° and II°. There is debate about the usefulness of root repair in knees with K&L grade III°. There is agreement that root repair should not be performed with K&L grade IV°, because PROMs and progression were superior and progression of OA is significantly less following repaired roots. In cases with a varus knee in patients with a degenerative posteromedial root tear, a high tibial osteotomy (HTO) to unload the medial compartment should be considered. By contrast, there is still debate about repairing the so-called ‘ramp lesion’ which occurs occasionally with ACL tears.

Fig. 6 One of the new solutions for complete meniscus replacement. (A) Schematic drawing of a free-floating, ring-shaped, polycarbonate-urethane implant (NuSurface by Active Implants®). (B) Arthroscopic photograph of the implant in place (the joint is filled with air in this photograph).

Biomechanical studies have proven the usefulness of a repair but there has been just one RCT so far showing no difference between leaving it alone or repairing it. The RCT is of lower quality and did not perform an a priori sample size calculation. Thus, it is unclear whether there was really no difference or whether the cohort was too small to find a difference.

Treatment of symptomatic cartilage lesions with chondral boundaries depends on the depths and the area of the lesion. Cartilage lesions > II° of ICRS classification are treated, lesions ≤ II° are left alone. Lesions of up to 25 mm² are treated with microfracturing or microdrilling, working best at the femoral condyle (Fig. 8). However, outcome improves up to two years and then it declines. One reason is that these techniques generate fibrocartilage repair with a low content of hyaline cartilage resulting in a variable volume of substitute cartilage. Furthermore, autologous chondrocyte transplantation (ACTs) have a higher failure rate after previous microfracture compared to ACT for primary lesions. Lesions above this size are better rather treated with ACT. ACT showed favourable results compared with microfracture after three years. However, ACT is not yet available in all countries. Furthermore, two surgeries are needed, and ACT is relatively expensive. Initially, cultured chondrocytes were transplanted into the defect, which was closed with a periosteum flap. Later on, a porcine type I/III collagen membrane was used to cover the cells fixed using fibrin glue or sutures during open surgery. The new generation can be applied arthroscopically by injecting a gel or applying small adhesive spheres which contain the cultured chondrocytes. A membrane to cover may not be necessary anymore. A further option to treat cartilage defects is osteochondral transplantation (OCT). It has shown good results for lesions up to 3 cm². However, after ten years, clinical results were inferior compared with ACT. Today it is mainly recommended for cartilage lesions affecting subchondral bone structures.
and that are limited in size. One should keep in mind that treating properly concomitant lesions such as in meniscus tears, patella instability, and leg malalignment will improve the outcome and the survival of the cartilage treatment.

For degenerative symptomatic cartilage lesions and osteoarthritis, conservative treatment plays a bigger role. Exercise, weight loss in overweight patients, cane use, bracing and NSAIDs were recommendations of the American College of Rheumatology/Arthritis Foundation. Disease-modifying drugs showed varied results. Intra-articular injections with corticosteroids (extended release), hyaluronic acid (high molecular weight) or platelet-rich plasma can help temporarily. Surgically, the above-mentioned

Fig. 7 Intra-operative picture of a knee in a figure-of-four position. Double-tunnel posterolateral root repair and an anatomical single bundle anterior cruciate ligament reconstruction (ACL-R) were performed. Tibial tunnels for ACL-R and posterolateral root repair were created independently to avoid compromising tensioning of the lateral meniscus.

Examples of Surgical Treatment of Cartilage Lesions

Fig. 8 Some examples of surgical treatment of cartilage lesions. Note. ACT, autologous chondrocyte transplantation; OCT, osteochondral transplantation.
biological techniques are also used in patients with degenerative cartilage lesions. However, there is low evidence. In general, these techniques need good quality surrounding cartilage, which is often not present in patients with degenerative cartilage lesions. Furthermore, patients are older, and with age biological repair capacity decreases and concomitant lesions, e.g. degenerative meniscus lesions, are more frequent, further compromising the outcome. Thus, using these biological techniques in patients with degenerative cartilage lesions is not the standard nowadays. Additionally, further techniques exist for degenerative cartilage lesions such as implanting a patient-specific metal plug into the chondral and bony defect. One of the most classical orthopaedic surgical techniques, correction of the leg axis to unload the affected compartment, is also one of the most effective. This is mostly done with a high tibial osteotomy (HTO). Nevertheless, more options exist, for example, the implantation of a spring on the medial side of the knee to unload the medial compartment. Biomechanically, the spring can lead to an unload of the medial compartment comparable to a 5° and 10° HTO. And of course, in severe OA or after failed prior OA surgeries, arthroplasty is an option, which will not be discussed in this article.

Conclusions

Cartilage and menisci have a close anatomical and functional relationship, which also plays a major role if one of them is damaged. Traumatic meniscal tears should be repaired to protect the articular cartilage and (partial) meniscus resection should be avoided. Degenerative meniscus lesions should be treated first conservatively, and only after failed conservative treatment (about three months) should arthroscopic partial meniscectomy be performed. Traumatic cartilage lesions should be treated depending on depth, affected area and possible bony pathology mainly by utilizing biological treatment options such as microfracture, OCT or autologous chondrocyte transplantation (ACT). For degenerative cartilage lesions, first-line treatment is conservative and second-line treatment is surgery with a variety of options, of which unloading the affected compartment is one of the most important.
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