Alternative and complementary therapies in osteoarthritis and cartilage repair

N. R. Fuggle · C. Cooper · R. O. C. Oreffo · A. J. Price · J. F. Kaux · E. Maheu · M. Cutolo · G. Honvo · P. G. Conaghan · F. Berenbaum · J. Branco · M. L. Brandi · B. Cortet · N. Veronese · A. A. Kurth · R. Matijevic · R. Roth · J. P. Pelletier · J. Martel-Pelletier · M. Vlaskovska · T. Thomas · W. F. Lems · N. Al-Daghri · O. Bruyère · R. Rizzoli · J. A. Kanis · J. Y. Reginster

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Abstract
Osteoarthritis (OA) is the most common joint condition and, with a burgeoning ageing population, is due to increase in prevalence. Beyond conventional medical and surgical interventions, there are an increasing number of ‘alternative’ therapies. These alternative therapies may have a limited evidence base and, for this reason, are often only afforded brief reference (or completely excluded) from current OA guidelines. Thus, the aim of this review was to synthesize the current evidence regarding autologous chondrocyte implantation (ACI), mesenchymal stem cell (MSC) therapy, platelet-rich plasma (PRP), vitamin D and other alternative therapies. The majority of studies were in knee OA or chondral defects. Matrix-assisted ACI has demonstrated exceedingly limited, symptomatic improvements in the treatment of cartilage defects of the knee and is not supported for the treatment of knee OA. There is some evidence to suggest symptomatic improvement with MSC injection in knee OA, with the suggestion of minimal structural improvement demonstrated on MRI and there are positive signals that PRP may also lead to symptomatic improvement, though variation in preparation makes inter-study comparison difficult. There is variability in findings with vitamin D supplementation in OA, and the only recommendation which can be made, at this time, is for replacement when vitamin D is deplete. Other alternative therapies reviewed have some evidence (though from small, poor-quality studies) to support improvement in symptoms and again there is often a wide variation in dosage and regimens. For all these therapeutic modalities, although controlled studies have been undertaken to evaluate effectiveness in OA, these have often been of small size, limited statistical power, uncertain blindness and using various methodologies. These deficiencies must leave the question as to whether they have been validated as effective therapies in OA (or chondral defects). The conclusions of this review are that all alternative interventions definitely require clinical trials with robust methodology, to assess their efficacy and safety in the treatment of OA beyond contextual and placebo effects.

Keywords Osteoarthritis · Cartilage · Alternative · Therapy · Treatment · Herbal

Introduction
Osteoarthritis (OA) is the most common form of arthritis and the global prevalence of knee OA alone is 3.8%, affecting over 250 million individuals worldwide [1]. OA is an increasingly major socioeconomic and public health issue, with the years lived with disability increasing by 64% from 1990 to 2010.

The current dogma is that OA may have differing causes but with a common, multi-tissue morphology including cartilage fibrillation, fissure and loss, subchondral bone changes and synovitis. OA is more prevalent in females than males and, although it can affect any joint, the most common anatomical sites include the knee, distal interphalangeal joints and hip [2]. Clinically, OA is characterized by joint pain, significant stiffness and leads to functional decline and a reduced quality of life for the affected individual.

There are a number of different treatments for OA including non-pharmacological and pharmacological approaches.
However, despite a number of well-written and well-considered guidelines [3–6], there is no direct advice regarding the application of what may be termed ‘alternative’ treatments including autologous chondrocyte implantation (ACI), autologous/heterologous mesenchymal stem cells (MSCs), platelet-rich plasma (PRP), vitamin D and other therapies (e.g. oral collagens, methylsulfonylmethane, curcumin, ginger). This lack of appropriate clinical advice and information is an issue for clinicians when considering how best to advise patients, especially as some of these therapies have a high profile in the lay press.

A current literature review was, therefore, performed and a working group of the European Society for Clinical and Economic Aspects of Osteoporosis and Osteoarthritis (ESCEO) was convened to review, evaluate and summarize current evidence regarding these putative OA treatments, and to provide expert opinion on their current role in the treatment of OA. We have classified the alternative therapies into surgical and medical approaches.

**Surgical therapy for cartilage loss**

Joint replacement is an established surgical technique focused on treating the end-stage of OA. For this reason, more minor surgical procedures have been developed to be used in the case of localized, traumatic or early disease with the aim of regenerating cartilage and rejuvenating the joint. In this section, we examine the evidence for the use of autologous stem cell and cartilage therapies as potential treatment options.

**Autologous chondrocyte implantation in knee cartilage defects**

ACI has a 30 year history [7] and is an established technique for the treatment of ulcerated cartilage and cartilage defects. It involves an initial cartilage biopsy, from which chondrocytes are cultured in vitro. In a second surgical procedure, a flap or membrane is then sutured (or glued) over the defect and the cultured chondrocytes are injected under this barrier. This process is summarized in Fig. 1. Over the last 10–15 years ACI has evolved (as bioengineering technology has improved) and now includes matrix-assisted ACI (MACI). The patient then undergoes very careful and graded rehabilitation to prevent the patch being dislodged.

Early randomised controlled trials (RCTs) data suggested no significant benefit of ACI when compared to the alternative surgical option of microfracture [8] and although a histologic improvement was observed [9], the clinical relevance of this is questionable. Indeed a Cochrane review in 2011 concluded that there was insufficient evidence to recommend the use of ACI [10].

The method developed over time to include collagen-covered ACI, and subsequently MACI, with the latter providing benefits including reduced size of the incision, greater surgical consistency, more consistent cell seeding, reduced periosteal hypertrophy and fewer adverse events [11–14].

Indeed, the matrix-applied method did perform significantly better than microfracture in the SUMMIT study.
(an RCT of 144 patients over 2 years) [15] in terms of clinical and functional outcomes. There was however no significant benefit over microfracture in MRI or histological outcomes. A key finding from these RCTs is that there was no correlation between the functional outcomes and evidence of structural repair when MRI is used, which is a potentially concerning finding.

ACI in combination with meniscal transplant allograft has good long-term outcomes with 75% still functioning well at 10-years (and 25% proceeding to arthroplasty). It is difficult to delineate whether the benefits of the procedure are due to ACI, meniscal transplant or indeed osteotomy (performed as part of the procedure) [16]. Similar results have been demonstrated in 57 patient with bipolar chondral lesions in the tibiofemoral compartment [17] with 75% having no radiographic progression at 10 years.

The cost of ACI and MACI are high, ranging from £4125 per patient to approximately £16,000 per MACI implant or £18,000 for a single vial of cells for ACI [18]. Therefore, in 2015 the technique was appraised by the National Institute for Health and Care Excellence (NICE) in the UK for the treatment of articular cartilage defects of the knee. The conclusions of this appraisal were that, while short-term clinical benefits were observed, the long-term clinical efficacy remained uncertain and the technique did not have robust evidence to demonstrate cost-effectiveness. Further research and evidence were recommended. The cost-effectiveness conclusions were considered harsh and were addressed in a consensus statement by UK knee surgeons, who drew attention to the estimated cost-effectiveness of ACI being between £7000 to £100,000 per Quality-Adjusted Life Year (QALY) (with the NICE threshold set at £20,000–30,000 per QALY).

There is a relative paucity of users, with, for example, only ten in the United Kingdom, which makes the development of large-scale research a challenge. MACI remains a potentially fruitful avenue for symptomatic therapy in early cartilage disease and traumatic cartilage lesions, though crucially not in OA.

Medical approaches

The scope of ‘non-surgical’ alternative therapies is large and, for this reason, this review focuses on the treatments which are likely to arise in clinical discussion with OA patients including autologous MSC injection, PRP, vitamin D and ‘other’ treatments.

Autologous mesenchymal stem cells

Articular cartilage is formed of a single cell type, the chondrocyte and a stable extracellular matrix that has no vascular, lymphatic or nervous supply. Subchondral bone provides mechanical and nutritional support and microfractures in this tissue can result in the release of undifferentiated mesenchymal stem cells (MSCs) from the bone marrow to facilitate the repair of chondral defects.

The repair capacity of MSCs has led to the development of techniques to directly inject MSCs locally into the joint following the ex vivo preparation of mesenchymal cells (Fig. 2). MSCs (also known as human or bone marrowstromal cells, multi-potent adult stem cells, mesenchymal progenitor cells and skeletal stem cells) have the ability to differentiate into the three tissue types, cartilage, bone and fat, and are invested, by definition, with an innate capacity for self-renewal and rapid proliferation. MSCs

Fig. 2 A depiction of the injectable and implantable options for delivery of MSCs and the potential sources of MSCs which are appropriate for each method

Variations in delivery mechanisms, MSC preparations and sources used: BM stromal cells, multi potential adult stem cells, human marrow stromal cells, or mesenchymal progenitors

- Free cell (modified)
- Expanded cell
- Source: adipose tissue, bone marrow, MSC, UC
- Expanded and activated cells

- Scaffold (artificial) + modified cells (1-step process)
  - Source: any

- Scaffold (artificial) + expanded and activated cells (2-step process)
  - Source: any

- Tissue engineered construct (1-step process)
  - Source: synovium, MSC
display paracrine anti-inflammatory and immunomodulatory properties [19, 20] and can be harvested from bone marrow (biopsy or aspirate) as well as from the stromal vascular fraction in adipose tissue. (Table 1) Autologous sources avoid any immunological concerns, however, there is concern that, as MSCs are cleared from the joint rapidly, the joint may simply be experiencing the benefits afforded by a ‘wash-out’.

The use of MSCs in OA is an area of burgeoning research and, at the time of writing, there were 182 studies recorded, in various states of progress and 3 systematic reviews which best address the use of stem cells in the treatment of OA [21–23].

The review performed by Hached and colleagues analysed a total of 44 trials of intra-articular injection of MSCs in the treatment of OA including bone marrow-derived, adipose tissue-derived and umbilical cord MSCs [21]. The review concluded that all three methods of acquisition of MSCs had evidence to support their use in the treatment of OA and that intra-articular injection of these cells was safe with very few side effects. An extensive review appraised 20 full-text records including systematic reviews, comprehensive reviews, clinical reviews and meta-analyses published between 2006 and 2016 which addressed the treatment of cartilage lesions with MSCs [23]. In their review, the authors noted that improvements in symptoms (pain and function) were more commonly reported than structural/tissue improvement. There was a low level of evidence for the intervention with a mere 25 items of Level I graded evidence and subsequently concluded that it was “unclear” if stem cells were an effective treatment for OA. Broadly, stem cell therapies were effective in symptomatic (pain) relief related to chondral defects and defects (or lesions) due to OA. The authors reported, overall, limited repair and integration with extensive variability in the results presented. The main issues recorded included significant variability in MSC sources, techniques for preparation, methods of administration and the range of co-interventions used (including micro-fracture, sub-chondral drilling, debridement, PRP).

There are very few studies which have demonstrated any degree of structural improvement in knee OA. Lam-Espinoza and colleagues [24], reported the results of an RCT of increasing doses (10 × 10⁶ or 100 × 10⁶) of bone marrow MSC, intra-articular injection against hyaluronic acid injection in 30 patients with OA (Kellgren–Lawrence grades II–IV). Participants were followed-up for 12 months and those in the MSC injection group had significant improvements in functionality and symptoms. Interestingly, only those in the high dose MSC injection group had statistically significant structural improvement in cartilage thickness on MRI at 12 months, opening the possibility of a dose–response. It should be noted that the inclusion of patients with such severe disease (Kellgren–Lawrence IV) suggests that the experimental group was substantially heterogeneous.

A pilot study by Orozco and colleagues [25] performed on patients with mild to severe knee OA (Kellgren–Lawrence grades II–IV) who received an intra-articular injection of (40 × 10⁶) bone marrow-derived MSCs, demonstrated improvement in pain, function and cartilage quality at 12 months. The pain relief was maintained at 2 years, while the objective cartilage improvement (on MRI) continued on a trajectory of improvement at 2 years [26]. However, it must be noted that this study only included 12 patients and so conclusions should be tentative at best.

Allogeneic MSC injection was demonstrated to be both feasible and safe, as reported in previous studies [27], though the observed negative outcomes include the generation of fibrocartilage, injection-related pain and swelling, infection post-bone marrow aspirate and a pulmonary embolus 2 weeks post-bone marrow aspirate [23]. In a further systematic review, only 2 serious adverse events (synovial effusion and unstable angina) were observed amidst 288 patients.

| MSCs Cell source | Advantages | Disadvantages |
|------------------|------------|--------------|
| Bone marrow (bone marrow conc. and bone marrow aspirate conc.) | High chondrogenic potential | High variability in MSC number |
| Adipose tissue and stromal vascular fraction | Relative ease of collection | MSC numbers and quality decline with age |
| Synovial membrane | Ease of harvest | MSC numbers decline with obesity |
| Umbilical cord MSCs | Large amount of tissue can be extracted | Lower chondrogenic potential |
| Peripheral blood, synovial fluid, amniotic fluid | Limited donor site morbidity | Limited number |
| | Major source of allogeneic cells | Chondrogenic potential variable |
| | Ease of harvest | |
| | Unlimited numbers | |
| | Relative ease of collection | High variability in MSC numbers |

Chondrogenic potential variable
Pers and colleagues studied three dosages of adipose tissue-derived MSCs (2 × 10^6, 10 × 10^6, 50 × 10^6 cells) in the Adipose Derived mesenchymal stromal cells in Patients with knee Osteoarthritis (ADIPOA) trial and found 2 × 10^6 was optimal in terms of functionality and pain relief at 9 months [28] and postulated MSCs may operate via innate and adaptive immune modulation [29]. This phase I trial is now in phase II (ADIPOA-2) and recruitment of 150 patients is underway.

Limitations and barriers to the routine application of MSCs for OA from this plethora of clinical studies were a consequence of (i) significant variation in MSC source, (ii) significant variation in MSC preparation protocols adopted, (iii) significant variation in MSC delivery approaches adopted and, (iv) significant variation in the number of different co-interventions with MSCs including micro-fracture, sub-chondral drilling, debridement, and PRP as well as hyaluronic acid, albumin and serum, osteophyte removal, and surgical interventions (ACL repair and high tibial osteotomy).

Nevertheless, the following factors were associated with increased efficacy of MSC injection for OA:

- Younger age
- Male gender
- Low BMI
- Small lesion/defect
- Early/mild to moderate OA severity

It should also be considered that these MSC treatments are currently not covered by many health insurance providers and the costs are high. For example, in the United States, the cost of a single stem-cell treatment for osteoarthritis was estimated at $5156 (95% CI $4550–5762) based on data from 273 centers [30].

In conclusion, the predominantly poor-quality, current literature suggests that symptoms, particularly pain, may improve with MSCs injection, however, evidence of structural improvement is unconvincing and positive effects appear to be observed in particular patient phenotypes. The overwhelming conclusion is a need to standardize the intervention if progress is to be made (Table 2). There is also a substantial need for phase II and III trials with the results of ADIPOA-2 being keenly awaited.

**Platelet-rich plasma**

Platelets play an important role in coagulation but also inflammation and PRP is a therapy which has been used extensively in equine tendinopathy [31] and has been investigated in the treatment of OA, particularly of the knee [32].

Platelet-rich plasma is a fluid which is rich in growth factors that stimulate cell proliferation, cellular migration, angiogenesis and the synthesis of the extracellular matrix including: platelet-derived growth factor (PDGF), tumor-like growth factor-β (TGF-β), fibroblast growth factor (FGF), vascular endothelial growth factor (VEGF), hepatocyte growth factor (HGF) and insulin-like growth factor-1 (IGF-1).

It is derived through centrifugation of a patient’s blood, with the aim of separating a plasma component which is rich in platelets (> 95% platelets) from whole blood which is poor in platelets (4% platelets). The PRP is then extracted and injected into the affected joint. The intricacies of preparation techniques vary and result in significantly different constituent cells (erythrocyte and leucocyte proportions), platelet concentrations and injection volumes [33] (Fig. 3). Indeed, there is a global schism in practice with Europeans preferring to use leukocyte-poor and Americans using leukocyte-rich PRP. PRP has been investigated in RCTs [34, 35] but the broad variation in preparation methods makes inter-trial comparison difficult and robust conclusions harder to ascertain and few are blinded. To emphasize this point we have synthesized and summarized some of the seminal studies below.

The issues surrounding the preparation of PRP are covered in a review of the techniques utilised in a number of RCTs and systematic reviews [36]. There is substantial

| Key factors for stem cell efficacy in OA                                                                 | Key factors limiting stem cell efficacy and clinical trial interpretation of MSC use in OA                                                                 |
|----------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| • Age (younger patients typically display better outcomes)                                               | • Significant variation in MSC source                                                                                                                                                           |
| • Gender (males typically better outcomes compared to female counterparts)                               | • Significant variation in MSC preparation protocols                                                                                                                                               |
| • BMI (lower BMI is associated with better outcomes)                                                     | • Variation in MSC Delivery                                                                                                                                                                     |
| • Lesion or defect size (better repair associated with smaller lesion size)                              | • Significant variation in the number of different co-interventions with MSCs                                                                                                                    |
| • Stage of OA (early OA, mild to moderate OA correlated with better outcomes)                           | Micro-fracture, sub-chondral drilling, debridement, and platelet rich plasma                                                                                                                     |
|                                                                                                        | Less common—hyaluronic acid, albumin and serum, osteophyte removal, and surgical interventions (ACL repair and high tibial osteotomy)                                                          |
variation in techniques including: the subject studied (severity of knee OA), PRP preparations, the inclusion of leukocytes, platelet count, number of injections delivered, interval/frequency of administration, volume of injection, whether fresh or freeze-thawed PRP were used, the use of anticoagulants and activating agents, separation techniques and any co-administered injections. With this in mind, a technical analysis was performed in 2017 to evaluate the similarities and differences between differing PRP formulations, in an attempt to determine the best preparation for the treatment of knee OA.

Filardo and colleagues [37, 38] performed a blinded trial in which they recruited participants with radiographic knee OA up to a Kellgren and Lawrence score of ≤ III, with 96 randomised to PRP and 96 to hyaluronic acid as a comparator. The PRP was centrifuged twice and PRP participants received 3 injections, once a week for three weeks and all participants were followed up for 12 months initially but extending to 5 years [39]. The key finding was that both treatments were equally effective in reducing knee OA symptoms and improving function over time but leucocyte-rich PRP was no more effective than hyaluronic acid.

To summarize the available evidence regarding PRP a number of systematic reviews have been performed [40–42]. PRP provided significant improvements in knee OA patient outcomes at 12 months and larger improvements were observed in those with milder radiographic disease (Kellgren and Lawrence ≤ II) [40].

Significant improvements in ‘patient recorded outcomes’ were also observed with PRP as opposed to hyaluronic acid at 3–6 months (WOMAC 28.5 vs. 43.4 respectively, \( p = 0.0008 \)) and 6–12 months (WOMAC 22.8 vs. 38.1, \( p = 0.0062 \)) [41].

A further systematic review published in 2018 (including 7 randomized placebo-controlled trials and 908 patients) sought to investigate the superiority of PRP over hyaluronic acid which was not demonstrated. In respect of PRP the minimal clinically important difference (MCID) was observed in 5 of the 7 papers, and suggested that differences in clinical outcomes could be due to variation in the preparation of PRP in terms of; centrifugation (speed, frequency, time-length, activating agents), administration (frequency, volume of injection) and post-administration rehabilitation protocols [42]. From a safety point of view, no local or systemic serious adverse events were noted in the reviewed articles.

Milants and colleagues used a previous definition of minimal clinically important improvement in pain (MCII) to determine whether an observed difference had any ‘meaningful’ effect in clinical practice. This was set at 15 out of 100 for absolute improvement and 20% for relative improvement for knee OA, as defined by Tubach et al. [43].

The Milants technical analysis included 19 RCTs, and studies were classified into two groups depending on outcomes with a ‘bad responder group’, defined as a response less than the minimal clinically important improvement (MCII) (\( n = 4 \) studies), and a ‘very good responder group’, defined as a response greater than twice the MCII (\( n = 7 \) studies). The reviewers contacted authors of the trials to obtain information regarding the preparation which was missing from the manuscript and PRP preparation was classified according to the Mishra (a classification in which PRP is divided into 4 types depending on 3 variables; white blood cells: increased or minimal, activation: yes or no, platelet content > 5 times patient baseline or ≤ 5 times patient baseline) and PAW (Platelet concentration, Activation prior to injection, White blood cell content).

In almost all studies with a very good responder group, PRPs were leukocyte-poor, activated prior to injection and platelets < 5 times baseline or between baseline and 750,000 platelets/µL), administered according to a lower number of injections (1 or 2 rather than 3), with a longer interval between injections (2 to 3 weeks per injection rather than once weekly) and a single (as opposed to double) spinning technique. The use of leukocyte-rich PRP was only found in the bad responder group. The use of calcium chloride and citrate was common in the very good responder group.

The cost of the PRP procedure is estimated at $714 (95% CI $691–737) based on data from 179 centers from across the United States [44].

In conclusion, although PRP may have repeated mild symptomatic benefits, there is yet to be experimentally robust demonstration of symptomatic and structural effects in the current literature. Research is required to better understand the mechanism of action, including investigation of the survival and location of platelet-derived factors within the joint following injection. In order for PRP to be considered...
within the dogma of recommended treatment for OA, at least one large, randomized, placebo-controlled trial and further investigation regarding preparation and dosage efficacy is required. This working group cannot, therefore, make a recommendation to use PRP as an intervention for OA.

Vitamin D

There is a secular trend toward decreased vitamin D levels, with serum concentrations averaging 49 ng/mL in the mid-twentieth century to approximately 23 ng/mL now, and with over a billion individuals being vitamin D deficient or insufficient [45]. Due to the role played by sunlight in the in vivo production of vitamin D, particularly low levels are observed at the extremes of latitude [46], and in winter months. Studies of seasonal gene expression have shown that some pro-inflammatory factors, including soluble IL-6 receptor and C-reactive protein have a peak expression in winter months and vitamin D receptor expression peaks in the summer months [47]. It is, therefore, interesting that certain diseases display similar seasonality and geography, including OA. This descriptive epidemiological observation is supported by basic scientific findings including [48]:

- There are receptors for vitamin D on chondrocytes which may play a role in the regulation of matrix metalloproteinases and prostaglandin E2 production
- Vitamin D stimulates proteoglycan synthesis in mature chondrocytes
- Vitamin D deficiency influences bone remodeling which may predispose to the development of OA

Despite these observations, four RCTs of vitamin D in OA have been performed in the United States (US) [49], India [50], the United Kingdom (UK) [51] and Australia [52]. None of these have demonstrated structural or symptomatic benefit in OA.

The pilot study performed in India [50] included 103 participants (59.4% females) with a baseline age of approximately 50 and a baseline 25-OH vitamin D of < 20 ng/mL. They found a significant reduction in knee pain and improvement in function but no significant alteration in radiographic knee OA at 12 months.

The placebo-controlled trial performed by McAlindon and colleagues [49] (in the US) included 146 female participants with a mean age at baseline of 62.4 years. At two years they found no improvements in knee symptoms, functional status or cartilage structure with vitamin D.

In the aforementioned UK placebo-controlled trial [51], despite an increase in serum 25-hydroxy-vitamin D (from approximately 20 to 30 µg/L) in the treatment group, no significant changes in symptomatic or radiographic knee OA were observed after 3 years in 474 participants (over the age of 50).

In Australia, Jin and colleagues demonstrated that in 209 patients with low vitamin D (12.5–60 nmol/L) treated for 2 years with monthly oral vitamin D3 (50,000 IU), there was no significant improvement in MRI-measured tibial cartilage volume or WOMAC knee pain score [52].

The relationship between vitamin D and knee OA has been investigated in a recent systematic review of 11 studies, which concluded that although vitamin D deficiency is associated with knee OA, the evidence regarding this association is inconsistent [53]. The studies included in the review were largely of cohort and cross-sectional design but also included two RCTs. The systematic review demonstrated that there was marked variation in the relationships between vitamin D and OA with a level of evidence (for an association of vitamin D deficiency with prevalent symptomatic knee OA) of ‘moderate’, while the relationship with prevalent radiographic knee OA was graded as ‘limited’. This negative conclusion supports that of a prior systematic review [54].

It should be acknowledged that vitamin D deficiency has been associated with a range of co-morbidities, of which OA is only one. However, the adverse effect profile of the supplement is favorable and should be strongly considered in those at risk of deficiency. Should there be systematic screening for vitamin D deficiency or systematic supplementation of vitamin D? This question is beyond the purview of this article and is dependent on many factors which are related to local healthcare systems and economic considerations.

We conclude by recommending that, when severe deficiency is diagnosed (especially in winter), vitamin D should be supplemented through the evidence that such supplementation ameliorates OA symptoms is inconclusive.

Other medical therapies

It should be noted that the medical therapies included in this section are very rarely mentioned in international guidelines, however, they are often the subject of discussions between patients and clinicians. Collagens, methylsulfonylmethane, S-adenosylmethionine, curcuma, harpagophytum and ginger are commonly used in the treatment of OA in many countries [55], with polyphenols, green tea, ‘Cat’s claw’ and dairy products also being mentioned.

Collagens: oral and intra-articular

Oral collagens are a rich source of amino acids, and, in OA, are purported to stimulate the joint to produce endogenous collagens in response to supplementation.

In 2016 a study investigated 190 patients, randomised to receive undenatured type II collagen (40 mg) or glucosamine
hydroxyshatinate and chondroitin sulfate or placebo [56]. The primary outcome was total WOMAC change from baseline with secondary outcomes being Lequesne index and pain VAS. After 6 months they found that undenatured type II collagen led to a significantly greater reduction in WOMAC compared to placebo (551 vs. 414, \( p = 0.002 \)) and compared to the glucosamine and chondroitin sulfate arm (551 vs. 454, \( p = 0.009 \)). In terms of secondary outcomes, there was a greater reduction in Lequesne index (2.9 vs. 2.1, \( p = 0.009 \)).

A further RCT investigated the performance of 5, once weekly 4 mL injections of polymerized collagen type I (of porcine origin) compared to sodium hyaluronate with assessments at 3 and 6 months [57]. The primary outcome was Lequesne index (measuring the severity of knee OA) at the 3 month time point with a visual analogue score for pain and SF-36 questionnaire also recorded. They found no significant differences between the groups for the above outcomes at either 3 or 6 months.

Gelatin, a form of collagen-rich in proline, was assessed in 52 patients as part of a randomised, placebo-controlled trial which found significant inter-group differences in several types of pain, however, an effect size was not reported, making extrapolation to clinical benefit difficult [58]. Undenatured collagen, an alternative form of collagen, was compared to glucosamine and chondroitin with 26 patients per group. This found no inter-group differences in the efficacy of the interventions [59].

In 2012, Van Vijven and colleagues published a systematic review of a variety of oral collagen at various doses comprised of 8 trials of collagen hydrolysates (3 versus placebo), gelatin (1 versus placebo) or undenatured collagen (versus glucosamine hydrochloride and chondroitin sulfate) [58]. The review concluded that there was ‘low’ grade evidence for the use of these in OA. Those treated with collagen hydrolysates included 313 treated patients (taking 10 g per day) against 297 on a placebo preparation and found a significant effect on symptoms (WOMAC pain (−0.48) with a significant but small effect size of 0.17) but no effect on the joint structure as assessed by MRI scan.

There are various preparations of oral or intra-articular collagens. Although widely used in a large number of countries, current data do not support a positive recommendation to treat OA patients despite a mild effect on symptoms (pain) and function.

**Methylsulfonylmethane (MSM)**

This dietary supplement is found in plants, fruits and vegetables, and can be taken alone or in combination with other supplements. There have been two, notable, placebo-controlled trials of MSM in patients with knee OA, both in approximately 50 patients over 12 weeks of follow-up. The first [60] involved a dosage schedule which resulted in 6 g per day and demonstrated a significant improvement in SF-36, WOMAC pain and function in the MSM group \( (p < 0.05) \). The second involved a dose of approximately 3 g per day demonstrated a significant improvement in WOMAC function but not in WOMAC pain or SF-36 compared to placebo [61].

A trial in knee OA randomised 118 patients between glucosamine, MSM, combination therapy or placebo and found a significant improvement in pain and Lequesne functional index at 12 weeks in all groups except the placebo arm.

A recent RCT examined the performance of glucosamine and chondroitin in combination with MSM versus glucosamine and chondroitin alone and versus placebo, in a population of 147 early knee OA patients (Kellgren and Lawrence grade I-II). This study demonstrated improvement in the groups which included MSM compared to the other treatment groups, in terms of pain VAS and WOMAC scores [62].

In conclusion, small trials did not demonstrate any major safety concerns for MSM treatment. Whether there is a symptomatic benefit over a short follow-up period is a question which would need to be answered through larger, well-designed trials and long term follow-up data are required.

**S-adenosylmethionine (SAMe)**

This is a substance produced from methionine in the liver. The treatment of hip and knee OA with SAMe was the subject of a Cochrane review of 4 randomised, placebo-controlled trials [63]. This review included 656 patients and demonstrated a significant improvement with SAMe compared to placebo but with a very minor effect on pain (4 mm on a 100 mm VAS) and function (2 mm on a 100 mm VAS), which are of questionable clinical significance. There was no significant difference in adverse effects or withdrawals but it should also be noted that the methodological and reporting quality were poor and that there was a moderate degree of inter-trial heterogeneity \( (I^2 = 54\%) \).

**Curcuma**

Curcuma (or curcumin) is an extract of turmeric, a yellow spice, and member of the zingiberaceae family. Both curcuma and ginger have roots in Ayurvedic and Chinese medicine [64] with curcuma manifesting an anti-inflammatory effect via cyclo-oxgenase (COX)-2, prostaglandins and leukotoxin inhibition. There is a wide variation in daily doses from 180 to 2000 mg, which makes direct, inter-study comparison problematic. Nevertheless, a meta-analysis was published in 2016 included 4 placebo-controlled trials in the context of knee OA; 2 trials versus ibuprofen and 1 versus diclofenac. In these trials of curcuma, improvements were seen in symptomatic measures and NSAID consumption (up
to 4 months) versus placebo but no significant superiority was noted versus ibuprofen or when added to diclofenac.

Although a few, idiosyncratic adverse effects were reported in the trials, the meta-analysis concluded that curcuma could be considered safe at daily doses of 4800 mg for 4 months [65].

A more recent meta-analysis, published in 2018, included 11 RCTs (N = 1009 patients) investigating the role of curminoids and boswellia (a gum-resin used in Ayurvedic medicine) in the treatment of knee OA [66]. There was some improvement in pain and function outcomes versus placebo, however, the conclusion was that evidence was currently too scant to allow the therapies to be included in clinical recommendations for treatment.

A trial of bio-optimized curcuma in the treatment of knee OA was reported in 2019 [67]. After 3 months of treatment, there were no statistical differences in intention to treat analyses for curcuma efficacy comparing the treatment to placebo arms for the co-primary endpoints (which are not included in those recommended by the European Medicines Agency or scientific societies). However, there was a significantly higher incidence of diarrhea in the curcuma group. In summary, experimental data regarding curcuma is sparse. The little evidence there supports a mild effect on OA symptoms. The sample sizes of the published trials are small, follow-up is short [65] and longer-term, robust studies are required before curcumin can be actively recommended from an efficacy stand-point.

**Harpagophytum**

Also known as ‘Devil’s claw’ Harpagophytum is an African plant which is thought to manifest anti-inflammatory effects, similar to curcuma, via inhibition of COX-2 and leukotxin. A systematic review by Gagnier and colleagues investigated the role of Harpagophytum in the treatment of lower back pain and OA, including 3 randomised, placebo-controlled trials of hip and knee OA (385 patients) [68]. This review concluded that there was ‘moderate’ evidence of effectiveness for the use of 60 mg of harpagophytum powder, though longer, higher-quality trials are required before it can be routinely recommended in clinical practice.

**Ginger**

Ginger, another member of the *Zingiberaceae* family, is thought to manifest anti-inflammatory effect via inhibition of COX, lipoxygenase, reduction in tumor necrosis factor and inflammatory prostaglandin production. There are data to suggest that ginger extract (Zintona EC®) is superior to placebo in terms of pain relief at 6 months (though efficacy was the same at 3 months) in a double-blind, randomised, placebo-controlled, crossover trial of knee OA [69].

Additionally, trials have demonstrated improvements in pain and mobility and reduced rescue medication usage intake versus placebo, though with more (mostly mild) gastrointestinal adverse events [70].

In 2015, the trial data were summarized and analyzed in a systematic review and meta-analysis of 5 randomised placebo-controlled trials, totaling 593 patients, which found a significant reduction in pain and disability with ginger. However, twice the rate of discontinuation was observed with ginger versus placebo (Relative Risk 2.33, 95% CI 1.04–5.22) [71]. A similar finding was reported in an earlier systematic review which noted “infrequent reports of mild, and predominantly gastrointestinal, adverse effects” [72].

**Conclusion**

In this review, we have synthesized the current evidence regarding alternative therapies for OA. Our findings are summarised in Table 3. Publication bias may be an issue with this group of treatments, however, this is not always the case (as seen throughout this review).

In summary, for all of the interventions covered in this review, issues of study design limit the degree to which inference can be made about clinical effectiveness in symptomatic OA. It is clear that none of these would currently clear the required hurdle for regulatory approval, were they to be assessed in like manner to current pharmaceutical interventions. However, there is also an insufficient basis for declaring them completely ineffective. They, therefore, remain an area in which further, appropriately designed, large, blinded, RCTs are an urgent necessity.
**Table 3**  A summary table of the key, take-home points for each of the interventions reviewed

| Alternative therapy                           | Key points                                                                 |
|-----------------------------------------------|---------------------------------------------------------------------------|
| Autologous chondrocyte implantation           | • Treatment for cartilage defects and not osteoarthritis                  |
|                                               | • Includes ACI and MACI                                                  |
|                                               | • Trial evidence to support symptomatic benefit                           |
|                                               | • Supported by NICE in the UK for specific patient group (including no previous knee surgery, limited evidence of knee osteoarthritis, large chondral defect) |
| Mesenchymal stem cell injection               | • There are multiple sources of MSCs with differing profiles of usage and limitations |
|                                               | • Multiple sources of MSCs lead to difficulties in directly comparing clinical trials |
|                                               | • Trial evidence to support symptomatic benefit                           |
|                                               | • Limited evidence to support structural benefit (MRI cartilage thickness) |
| Platelet-rich plasma                          | • Trial evidence to support symptomatic benefit                           |
|                                               | • Multiple methods of preparation lead to difficulties in directly comparing clinical trials |
| Vitamin D                                     | • Evidence of efficacy in clinical trials is equivocal                    |
|                                               | • Recommendation to provide supplementation to those patients with evidence of depleted levels of 25OH-vitamin D |
| Other alternative therapies                   | • Very limited clinical trial data to suggest the efficacy of oral collagens, MSM, SAMe, curcuma and ginger |
|                                               | • Adverse events: largely rare, though ginger appears to be associated with an increased risk of mild gastro-intestinal adverse events |

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References

1. Cross M, Smith E, Hoy D et al (2014) The global burden of hip and knee osteoarthritis: estimates from the global burden of disease 2010 study. Ann Rheum Dis 73:1323–1330. https://doi.org/10.1136/annrheumdis-2013-204763

2. Van Saase JL, Van Romunde LK, Cats A et al (1989) Epidemiology of osteoarthritis: Zoetermeer survey. Comparison of radiological osteoarthritis in a Dutch population with that in 10 other populations. Ann Rheum Dis 48:271–280

3. McAlindon TE, Bannuru RR, Sullivan MC et al (2014) OARSI guidelines for the non-surgical management of knee osteoarthritis. Osteoarthritis Cartilage 22:363–388. https://doi.org/10.1016/j.joca.2014.01.003

4. Bruyere O, Cooper C, Pelletier JP et al (2014) An algorithm recommendation for the management of knee osteoarthritis in Europe and internationally: a report from a task force of the European Society for Clinical and Economic Aspects of Osteoporosis and Osteoarthritis (ESCEO). Semin Arthritis Rheum 44:253–263. https://doi.org/10.1016/j.semarthritis.2014.05.014

5. Hochberg MC, Altman RD, April KT et al (2012) American College of Rheumatology 2012 recommendations for the use of non-pharmacologic and pharmacologic therapies in osteoarthritis of the hand, hip, and knee. Arthritis Care Res (Hoboken) 64:465–474

6. Jordan KM, Arden NK, Doherty M et al (2003) EULAR Recommendations 2003: an evidence based approach to the management of knee osteoarthritis: Report of a Task Force of the Standing Committee for International Clinical Studies Including Therapeutic Trials (EULAR). Ann Rheum Dis 62:1145–1155

7. Brittberg M, Lindahl A, Nilsson A et al (1994) Treatment of deep cartilage defects in the knee with autologous chondrocyte transplantation. N Engl J Med 331:889–895. https://doi.org/10.1056/nejm199410063311401

8. Knutsen G, Drogset JO, Engerbretsen L et al (2007) A randomized trial comparing autologous chondrocyte implantation with microfracture. Findings at five years. J Bone Joint Surg Am 89:2105–2112. https://doi.org/10.2106/jbjs.g.00003

9. Saris DB, Vanlauwe J, Victor J et al (2008) 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 36:235–246. https://doi.org/10.1177/0363546507311095

10. Vasiiliadis HS, Wasiaik J (2010) Autologous chondrocyte implantation for full thickness articular cartilage defects of the knee. Cochrane Database Syst Rev 10:CD003323

11. Sohn DH, Lottman LM, Lum LY et al (2002) Effect of gravity on localization of chondrocytes implanted in cartilage defects. Clin Orthop Related Res 394:254–262

12. Brittberg M (2010) Cell carriers as the next generation of cell therapy for cartilage repair: a review of the matrix-induced autologous chondrocyte implantation procedure. Am J Sports Med 38:1259–1271

13. Marlovits S, Aldrian S, Wondrasch B et al (2012) Clinical and radiological outcomes 5 years after matrix-induced autologous chondrocyte implantation in patients with symptomatic, traumatic chondral defects. Am J Sports Med 40:2273–2280

14. Ebert JR, Robertson WB, Lloyd DG et al (2010) A prospective, randomized comparison of traditional and accelerated approaches to postoperative rehabilitation following autologous chondrocyte implantation: 2-year clinical outcomes. Cartilage 1:180–187

15. Saris D, Price A, Widuchowski W et al (2014) Matrix-applied characterized autologous cultured chondrocytes versus microfracture: two-year follow-up of a prospective randomized trial. Am J Sports Med 42:1384–1394. https://doi.org/10.1177/036354514528093

16. Ogura T, Bryant T, Minas T (2016) Biological knee reconstruction with concomitant autologous chondrocyte implantation and meniscal allograft transplantation: mid- to long-term outcomes. Orthop J Sports Med 4:2325967116668490. https://doi.org/10.1177/2325967116668490

17. Ogura T, Bryant T, Mosier BA et al (2018) Autologous chondrocyte implantation for bipolar chondral lesions in the tibiofemoral compartment. Am J Sports Med 46:1371–1381. https://doi.org/10.1177/0363545518756977

18. NICE (2015) Autologous chondrocyte implantation for repairing symptomatic articular cartilage defects of the knee. National Institute for Health and Care Excellence. London

19. Harrell CR, Markovic BS, Fellbaum C et al (2019) Mesenchymal stem cell-based therapy of osteoarthritis: Current knowledge and future perspectives. Biomed Pharmacother 109:2318–2326. https://doi.org/10.1016/j.biopha.2018.11.099

20. Pastides P, Chimutengwende-Gordon M, Maffulli N et al (2013) Stem cell therapy for human cartilage defects: a systematic review. Osteoarthr Cartil 21:646–654. https://doi.org/10.1016/j.joca.2013.02.008

21. Hached F, Vinatier C, Le Visage C et al (2017) Biomaterial-assisted cell therapy in osteoarthritis: From mesenchymal stem cells to cell encapsulation. Best Pract Res Clin Rheumatol 31:730–745. https://doi.org/10.1016/j.berh.2018.05.002

22. Jevotovsky DS, Alfonso AR, Einhorn TA et al (2018) Osteoarthritis and stem cell therapy in humans: a systematic review. Osteoarthritis Cartilage 26:711–729. https://doi.org/10.1016.j.joca.2018.02.906

23. Hart D (2017) Mesenchymal Stem Cells: The hope, the hype and the reality in the treatment of osteoarthritis, A knowledge synthesis of clinical research (2010-2016) emphasizing the safety and efficacy of stem cell treatment for osteoarthritis. Bone and joint health strategic clinical network (BJH SCN)

24. Lamo-Espinosa JM, Mora G, Blanco JF et al (2016) Intra-articular injection of two different doses of autologous bone marrow mesenchymal stem cells versus hyaluronic acid in the treatment of knee osteoarthritis: multicenter randomized controlled clinical trial (phase I/II). J Transl Med 14:246. https://doi.org/10.1186/s12967-016-0998-2

25. Orozco L, Munar A, Soler R et al (2013) Treatment of knee osteoarthritis with autologous mesenchymal stem cells: a pilot study. Transplantation 95:1535–1541. https://doi.org/10.1097/TP.0b013e318291a2da2a

26. Orozco L, Munar A, Soler R et al (2014) Treatment of knee osteoarthritis with autologous mesenchymal stem cells: two-year follow-up results. Transplantation 97:e318291a2da2a

27. Vega A, Martin-Ferrero MA, Del Canto F et al (2015) Treatment of knee osteoarthritis with allogeneic bone marrow mesenchymal stem cells: a randomized controlled trial. Transplantation 99:1681–1690. https://doi.org/10.1097/TP.0000000000000678

28. Pers YM, Rackwitz L, Ferreira R et al (2018) Adipose mesenchymal stromal cell-based therapy for severe osteoarthritis of the knee: a phase I dose-escalation trial. Stem Cells Transl Med 7:857–866. https://doi.org/10.1002/sctm.2015-0245

29. Pers YM, Quentin J, Feirreira R et al (2018) Injection of adipose-derived stromal cells in the knee of patients with severe osteoarthritis has a systemic effect and promotes an anti-inflammatory phenotype of circulating immune cells. Theranostics 8:5519–5528. https://doi.org/10.7150/thno.27674

30. Puzzi NS, Ng M, Chuhati M et al (2018) The stem-cell market for the treatment of knee osteoarthritis: a patient perspective. J Knee Surg 31:551–556. https://doi.org/10.1055/s-0037-1604443

31. Ortved KF (2018) Regenerative medicine and rehabilitation for tendinous and ligamentous injuries in sport horses. Vet Clin Aging Clinical and Experimental Research (2020) 32:547–560
N Am Equine Pract 34:359–373. https://doi.org/10.1016/j.jcnev.2018.04.012

32. Mascaréñas R, Saltzman BM, Fortier LA et al (2015) Role of platelet-rich plasma in articular cartilage injury and disease. J Knee Surg 28:3–10. https://doi.org/10.1055/s-0034-1384672

33. Kauth JF, Le Goiff C, Seidel L et al (2011) Comparative study of five techniques of preparation of platelet-rich plasma. Pathol Biol (Paris) 59:157–160. https://doi.org/10.1016/j.pathbio.2009.04.007

34. Lin KY, Yang CC, Hsu CJ et al (2019) Intra-articular Injection of platelet-rich plasma is superior to hyaluronic acid or saline solution in the treatment of mild to moderate knee osteoarthritis: a randomized, double-blind, triple-placebo, placebo-controlled clinical trial. Arthroscopy 35:106–117. https://doi.org/10.1016/j.arthro.2018.06.035

35. Huang Y, Liu X, Xu X et al (2019) Intra-articular injections of platelet-rich plasma, hyaluronic acid or corticosteroids for knee osteoarthritis: a prospective randomized controlled study. Orthopade 48:239–247. https://doi.org/10.1007/s00132-018-03659-5

36. Milants C, Bruyère O, Kaux J-F (2017) Responders to platelet-rich plasma in osteoarthritis: a technical analysis. BioMed Res Int 2017:7538604. https://doi.org/10.1155/2017/7538604

37. Filardo G, Kon E, Di Martino A et al (2012) Platelet-rich plasma vs hyaluronic acid to treat knee degenerative pathology: study design and preliminary results of a randomized controlled trial. BMC Musculoskelet Disord 13:229. https://doi.org/10.1186/1471-2474-13-229

38. Filardo G, Di Matteo B, Di Martino A et al (2015) Platelet-rich plasma intra-articular knee injections show no superiority versus viscosupplementation: a randomized controlled trial. Am J Sports Med 43:1575–1582. https://doi.org/10.1177/0363546515582027

39. Di Martino A, Di Matteo B, Papio T et al (2018) Platelet-rich plasma versus hyaluronic acid injections for the treatment of knee osteoarthritis: results at 5 years of a double-blind, randomized controlled trial. Am J Sports Med. https://doi.org/10.1177/0363546518465188

40. Campbell KA, Saltzman BM, Mascaréñas R et al (2015) Does intra-articular platelet-rich plasma injection provide clinically superior outcomes compared with other therapies in the treatment of knee osteoarthritis? A systematic review of overlapping meta-analyses. Arthroscopy 31:2213–2221. https://doi.org/10.1016/j.arthro.2015.03.041

41. Meheux CJ, McCulloch PC, Lintner DM et al (2016) Efficacy of intra-articular platelet-rich plasma injections in knee osteoarthritis: a systematic review. Arthroscopy 32:495–505. https://doi.org/10.1016/j.arthro.2015.08.005

42. Di Y, Han C, Zhao L et al (2018) Is local platelet-rich plasma injection clinically superior to hyaluronic acid for the treatment of knee osteoarthritis? A systematic review of randomized controlled trials. Arthritis Res Ther 20:128. https://doi.org/10.1186/s13075-018-1621-0

43. Tubach F, Ravaud P, Martin-Mola E et al (2012) Minimum clinically important improvement and patient acceptable symptom state in pain and function in rheumatoid arthritis, ankylosing spondylitis, chronic back pain, hand osteoarthritis, and hip and knee osteoarthritis: Results from a prospective multinational study. Arthritis Care Res (Hoboken) 64:1699–1707. https://doi.org/10.1002/acr.21747

44. Puzzi NS, Ng M, Kantor A et al (2019) What is the price and claimed efficacy of platelet-rich plasma injections for the treatment of knee osteoarthritis in the United States? J Knee Surg 32:879–885. https://doi.org/10.1016/j.jknee.2018.03.038

45. Barger-Lux MJ, Heaney RP (2002) Effects of average summer sun exposure on serum 25-hydroxyvitamin D and calcium absorption. J Clin Endocrinol Metab 87:4952–4956. https://doi.org/10.1210/jc.2002-020636

46. Koch T (2014) 1831: the map that launched the idea of global health. Int J Epidemiol 43:1014–1020. https://doi.org/10.1093/ije/dyu099

47. Dopico XC, Evangelou M, Ferreira RC et al (2015) Widespread seasonal gene expression reveals annual differences in human immunity and physiology. Nat Commun 6:7000. https://doi.org/10.1038/ncomms8000

48. Tetlow LC, Woolley DE (2001) Expression of vitamin D receptors and matrix metalloproteinases in osteoarthritic cartilage and human articular chondrocytes in vitro. Osteoarthr Cartil 9:423–431. https://doi.org/10.1010/joca.2000.0408

49. McAlindon T, LaValley M, Schneider E et al (2013) Effect of vitamin D supplementation on progression of knee pain and cartilage volume loss in patients with symptomatic osteoarthritis: a randomized controlled trial. JAMA 309:155–162. https://doi.org/10.1001/jama.2012.164487

50. Sanghi D, Mishra A, Sharma AC et al (2013) Does vitamin D improve osteoarthritis of the knee: a randomized controlled pilot trial. Clin Orthop Relat Res 471:3556–3562. https://doi.org/10.1007/s11999-013-3201-6

51. Arden NK, Cro S, Sheard S et al (2016) The effect of vitamin D supplementation on knee osteoarthritis, the VIDEO study: a randomised controlled trial. Osteoarthr Cartil 24:1858–1866. https://doi.org/10.1016/j.joca.2016.05.020

52. Jin X, Jones G, Cicuttini FM et al (2016) Effect of vitamin D supplementation on tibial cartilage volume and knee pain among patients with symptomatic knee osteoarthritis: a randomized clinical trial. JAMA 315:1005–1013. https://doi.org/10.1001/jama.2016.1961

53. Vaishya R, Vijay V, Lama P et al (2019) Does vitamin D deficiency influence the incidence and progression of knee osteoarthritis? A literature review. J Clin Orthop Trauma 10:9–15. https://doi.org/10.1016/j.jcot.2018.05.012

54. Diao N, Yang B, Yu F (2017) Effect of vitamin D supplementation on knee osteoarthritis: a systematic review and meta-analysis of randomized clinical trials. Clin Biochem 50:1312–1316. https://doi.org/10.1016/j.clinbiochem.2017.09.001

55. Lim YZ, Hussain SM, Cicuttini FM et al (2019) Bioactive foods as dietary interventions for arthritis and related inflammatory diseases. Int: Watson RR, Preedy VR (eds) Bioactive foods in chronic disease states. Academic Press, Boston, pp 97–137. https://doi.org/10.1016/B978-0-12-813820-5.00006-4

56. Lugo JP, Saiyed ZM, Lane NE (2016) Efficacy and tolerability of an undenatured type II collagen supplement in modulating knee osteoarthritis symptoms: a multicenter randomized, double-blind, placebo-controlled study. Nutr J 15:14. https://doi.org/10.1186/s12937-016-0130-8

57. Martin Martin LS, Massafa U, Bizzi E et al (2016) A double blind randomized active-controlled clinical trial on the intra-articular use of Md-Knee versus sodium hyaluronate in patients with knee osteoarthritis (“Joint”). BMC Musculoskelet Disord 17:94. https://doi.org/10.1186/s12891-016-0948-4

58. Van Vijven JP, Luijsterburg PA, Verhagen AP et al (2012) Symptomatic and chondroprotective treatment with collagen derivatives in osteoarthritis: a systematic review. Osteoarthr Cartil 20:809–821. https://doi.org/10.1016/j.joca.2012.04.008

59. Crowley DC, Lau FC, Sharma P et al (2009) Safety and efficacy of undenatured type II collagen in the treatment of osteoarthritis of the knee: a clinical trial. Int J Med Sci 6:312–321

60. Kim LS, Axelrod LJ, Howard P et al (2006) Efficacy of methylsulfonylmethane (MSM) in osteoarthritis pain of the knee: a pilot clinical trial. Osteoarthr Cartil 14:286–294. https://doi.org/10.1016/j.joca.2005.10.003

61. Debi EM, Agar G, Fichman G et al (2011) Efficacy of methylsulfonylmethane supplementation on osteoarthritis of the knee: a randomized controlled study. BMC Complement Altern Med 11:50. https://doi.org/10.1186/1472-6882-11-50
62. Labis AMT, Siagian C, Wonggosummana E et al (2017) Comparison of Glucosamine-Chondroitin Sulfate with and without Methylsulfonylmethane in Grade I-II Knee Osteoarthritis: A Double Blind Randomized Controlled Trial. Acta Med Indones 49:105–111

63. Rutjes AW, Nuesch E, Reichenbach S et al (2009) S-Adenosylmethionine for osteoarthritis of the knee or hip. Cochrane Database Syst Rev 4:CD007321. https://doi.org/10.1002/14651858.CD007321.pub3

64. Goel A, Kunnumakkara AB, Aggarwal BB (2008) Curcumin as “Curecumin”: from kitchen to clinic. Biochem Pharmacol 75:787–809. https://doi.org/10.1016/j.bcp.2007.08.016

65. Daily JW, Yang M, Park S (2016) Efficacy of turmeric extracts and curcumin for alleviating the symptoms of joint arthritis: a systematic review and meta-analysis of randomized clinical trials. J Med Food 19:717–729. https://doi.org/10.1089/jmf.2016.3705

66. Bannuru RR, Osani MC, Al-Eid F et al (2018) Efficacy of curcumin and Boswellia for knee osteoarthritis: systematic review and meta-analysis. Semin Arthritis Rheum 48:416–429. https://doi.org/10.1016/j.semarthrit.2018.03.001

67. Henrotin Y, Malaise M, Wittoek R et al (2019) Bio-optimized Curcuma longa extract is efficient on knee osteoarthritis pain: a double-blind multicenter randomized placebo controlled three-arm study. Arthritis Res Ther 21:179. https://doi.org/10.1186/s13075-019-1960-5

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Affiliations

N. R. Fuggle1 · C. Cooper1,2 · R. O. C. Oreffo3 · A. J. Price4 · J. F. Kaux5 · E. Maheu5 · M. Cutolo6 · G. Honvo8 · P. G. Conaghan9 · F. Berenbaum10 · J. Branco11,12 · M. L. Brandi13 · B. Cortet14 · N. Veronese15 · A. A. Kurth16 · R. Matijevic17 · R. Roth18 · J. P. Pelletier19 · J. Martel-Pelletier19 · M. Vlaskovska20 · T. Thomas21,22 · W. F. Lems23 · N. Al-Daghri24 · O. Bruyère8 · R. Rizzoli25 · J. A. Kanis26,27 · J. Y. Reginster8,24,28

1 MRC Lifecourse Epidemiology Unit, University of Southampton, Tremona Road, Southampton SO16 6YD, UK
2 NIHR Musculoskeletal Biomedical Research Unit, University of Oxford, Oxford, UK
3 Bone and Joint Research Group, Centre for Human Development, Stem Cells and Regeneration, Institute of Developmental Sciences, University of Southampton, Southampton, UK
4 Nuffield Department of Orthopaedics, Rheumatology and Musculoskeletal Sciences, University of Oxford, Oxford, UK
5 Department of Physical and Rehabilitation Medicine & Sports Traumatology, FIFA Medical Centre of Excellence, IOC Research Centre for Prevention of Injury and Protection of Athlete Health, FIMS Collaborative Center of Sports Medicine, University Hospital and University of Liège, Liege, Belgium
6 Rheumatology Department, AP-HP, Saint-Antoine Hospital, 4 Blvd. Beaumarchais, Paris, France
7 Research Laboratory and Academic Division of Clinical Rheumatology, Department of Internal Medicine, University of Genoa, Genoa, Italy
8 WHO Collaborating Centre for Public Health Aspects of Musculoskeletal Health and Aging, Liege, Belgium
9 Leeds Institute of Rheumatic and Musculoskeletal Medicine, University of Leeds & NIHR Leeds Biomedical Research Centre, Leeds, UK
10 Department of Rheumatology, Sorbonne Université, INSERM CRSA, AP-HP Saint-Antoine Hospital, Paris, France
11 Centro Hospitalar de Lisboa Ocidental- Hospital Egas Moniz, Lisbon, Portugal
12 CEDOC / NOVA Medical School, Nova University of Lisbon, Lisbon, Portugal
13 Department of Surgery and Translational Medicine, University of Florence, Florence, Italy
14 Department of Rheumatology and EA 4490, Lille University Hospital, Lille, France
15 National Research Council, Neuroscience Institute, Aging Branch, Padua, Italy
16 Department of Orthopaedic Surgery, Themistocles Gluck Hospital, Ratingen, Germany
17 Faculty of Medicine, Clinic for Orthopedic Surgery and Traumatology, Clinical Center of Vojvodina, University of Novi Sad, Novi Sad, Serbia
18 Institute of Outdoor Sports and Environmental Science, German Sport University, Cologne, Germany
19 Osteoarthritis Research Unit, University of Montreal Hospital Research Centre (CRCHUM), Montreal, QC, Canada

20 Medical Faculty, Department of Pharmacology, Medical University Sofia, 2, Zdrave Str, 1431 Sofia, Bulgaria

21 Department of Rheumatology, Hôpital Nord, CHU de Saint-Étienne, Saint-Étienne, France

22 INSERM U1059, Université de Lyon, Saint-Étienne, France

23 Location VU Medical Center, Department of Rheumatology and Immunology, Amsterdam University Medical Center, Amsterdam, The Netherlands

24 Chair for Biomarkers Research, Biochemistry Department, College of Science, King Saud University, Riyadh, Kingdom of Saudi Arabia

25 Division of Bone Diseases, Geneva University Hospitals and Faculty of Medicine, Geneva, Switzerland

26 Mary McKillop Health Institute, Australian Catholic University, Melbourne, Australia

27 Centre for Metabolic Bone Diseases, University of Sheffield Medical School, Sheffield, UK

28 Department of Public Health, Epidemiology and Health Economics, University of Liège, CHU Sart Tilman B23, 4000 Liège, Belgium