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Evaluation of symptomatic slow-acting drugs in osteoarthritis using the GRADE system

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Results: Chondroitin sulfate, diacereine, glucosamine sulfate, avocado/soybean unsaponifiables and hyaluronic acid have demonstrated pain reduction and physical function improvement with very low toxicity, with moderate to high quality evidence. Even if pre-clinical data and some preliminary in vivo studies have suggested that oral calcitonin and strontium ranelate could be of potential interest in OA, additional well-designed studies are needed.

Conclusion: In the benefit/risk ratio, the use of chondroitin sulfate, diacereine, glucosamine sulfate, avocado/soybean unsaponifiables and hyaluronic acid could be of potential interest for the symptomatic management of OA.

Background

Osteoarthritis (OA) is a progressive disorder characterized by destruction of articular cartilage and subchondral bone associated with synovial changes [1,2]. This degenerative condition affects aging men and women [3]. The two most affected location for pain and physical disability in adults are hip and knee [4]. Because of its important prevalence worldwide, OA represents a huge burden in terms of individual, as well as public health resources utilization [5]. Pharmacological and non-pharmacological proce-
dures have demonstrated their efficacy to stop or decrease progression of this condition. Among pharmacological treatments, symptomatic slow-acting drugs have been largely studied over the last decade.

Most countries face common challenges in delivering consistent, appropriate and high quality health care standards within the limits of available resources. Clinical guidelines are one of the most important options to support and promote good clinical practices, and subsequently to make patient care more effective and efficient. However, to ensure that clinical guidelines actually meet this objective, they should follow a strict, validated methodology.

Over the last decade, several scientific societies involved in OA produced guidelines for the management of hip, knee and hand to improve quality and effectiveness of patients care [6-11]. Nevertheless, the most recent versions which have been prepared by prestigious institutions, such as the American College of Rheumatology (ACR) or the European League Against Rheumatism (EULAR) do not include the latest original research papers [12-28]. Furthermore, the recommendations established on the same topic by these two groups often differ. Part of the reasons are the lack of evidence, different interpretation of evidence, unsystematic guideline development methods, influence of professional bodies, cultural and socio-economic factors and differences in health care systems [29-31].

For all these reasons the European Society for Clinical and Economic Aspects of Osteoporosis and Osteoarthritis (ESCEO) experts found appropriate to build their own recommendations for the use of symptomatic slow-acting drugs in osteoarthritis (SYSADOA) based on the GRADE system [32,33]. The acronym of GRADE stands for Grading of Recommendations Assessment, Development and Evaluation.

The GRADE system is based on a sequential assessment of the quality of evidence, followed by assessment of the balance between benefits versus downsides and subsequent judgment about the strength of recommendations. Because frontline consumers of recommendations will be most interested in the best course of action, the GRADE system places the strength of the recommendation first, followed by the quality of the evidence. Separating the judgments regarding the quality of evidence from judgments about the strength of recommendations is a critical and specific feature of this new grading system.

GRADE has two level of recommendation: strong and weak (Table 1). As a matter of fact, recommendations to administer, or not administer, an intervention, should be based on trade-offs between benefits and risks, burden and, where possible, costs. If benefits outweigh risks and burden, experts will recommend that clinicians offer a treatment to patients sustaining typical symptoms of the disease. The uncertainty associated with the trade-off between the benefits and risks and burdens will determine the strength of recommendations.

### Methods

A systematic and exhaustive search of the meta-analysis and randomized controlled trials published from 1950 until December 2007 has been undertaken, using several tools, such as Medline, Old Medline, Embase, CINAHL, Science Citation Index through Web of Science, Allied Complementary Medicine and Cochrane Library databases. The search in the Cochrane Library included the Cochrane Reviews, Abstracts of Quality Assessed Systematic Reviews, The Cochrane Controlled Trial Register, NHS Economic Evaluation Database, Health Technology Assessment Database and NHS Economic Evaluation Bibliography Details Only.

### Table 1: Strength of guideline recommendations, consensus-based statements, and implication to quality of evidence

<table>
<thead>
<tr>
<th>Recommendation or statement</th>
<th>Description in GRADE approach</th>
<th>Interpretation</th>
</tr>
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</table>
| Strong guideline recommendation | We recommend (should) | 1. Most individuals should receive the intervention, assuming that they have been informed about and have understood its benefits, harms and burden.  
2. Most individuals would want the recommended course of action and only a small proportion would not.  
3. The recommendation could unequivocally be used for policy making. |
| Weak guideline recommendation | We suggest (might) | 1. The majority of individuals would want to suggested course of action, but an appreciable proportion would not.  
2. Values and preferences vary widely.  
3. Policy making will require extensive debates and involvement of many stakeholders. |
Quality of the evidence has been assessed using the grade four-category system (high, moderate, low and very low quality) (Table 2).

Factors that are considered in classifying evidence are: the study design and rigour of its execution, the consistency of results and how well the evidence can be directly applied to patients, interventions, outcomes and comparator. Other important factors are whether the data are sparse or imprecise and whether there is potential for reporting bias. Using this approach, assessments of the quality of evidence for each important outcome take into account the study design, limitations of the studies, consistency of the evidence across studies, the directness of the evidence, and the precision of the estimate.

Obviously, all outcomes (e.g. pain, function, NSAIDS consumption, carry-over effect, harm, global patient satisfaction, use of walking aids, structure modification, and evaluation by the GP) have not the same importance. The importance of each outcome has then been scored (10 mm VAS) independently by each expert. Only outcomes considered as important (VAS > 6 mm) by all experts were discussed in these recommendations.

For each intervention considered, the panel formulated a consensus recommendation based on the panel members' judgments regarding the balance between the benefits, harms (adverse effects), burdens (e.g., taking medication daily), costs, and values and preferences (the desirability or preference that individuals exhibit for a particular outcome) of the intervention. Source of funding has not been considered. Then, recommendations have been classified as "strong" or "weak."

**Results**

The experts considered that pain, function and harm are of primary interest in the evaluation of SYSADOA.

The following interventions were taken into consideration:

- Avocado/soybean unsaponifiables [34-36].
- Chondroitin sulfate [37-40].
- Diacereine [41,42].
- Glucosamine sulfate [38,39,43-45].
- Hyaluronic acid [46-52].
- Oral calcitonin [53].
- Risedronate [54,55].
- Strontium ranelate [56].

Following the GRADE system, the study design for all trials included in the review of evidence for chondroitin sulfate, diacereine, glucosamine sulfate, hyaluronic acid and risedronate was randomised controlled trial which is scored as a high type of evidence. As requested from the methodology of GRADE, study quality was also assessed by reviewing whether the studies had limitations or flaws. The following limitations were noted, leading frequently to a decrease in the quality of evidence: methods of randomisation were not clearly reported, allocation concealment was not reported or unclear, some trials were single-blinded and frequently the method of blinding was not reported in detail, incomplete descriptions of withdrawals and dropouts were reported, analyses were based on the per protocol or completor population and not on the intention-to-treat population, heterogeneity not tested in meta-analysis, large heterogeneity between meta-analyses statistically significant differences were reported at baseline between treatment and control groups, different severity of knee OA patients were included, large results discrepancies (e.g. between studies or between meta-analysis and large recent studies), nature of the placebo comparator, very few number of patient included, post-hoc analysis.

Quality evidence could also decrease when considering meta-analysis (e.g. moderate and high heterogeneity between RCT, or heterogeneity between meta-analyses).

<table>
<thead>
<tr>
<th>Grade</th>
<th>Definition</th>
<th>Underlying Methodology</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>Further research is very unlikely to change our confidence in the estimate of effect</td>
<td>RCT or meta-analysis</td>
</tr>
<tr>
<td>Moderate</td>
<td>Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate</td>
<td>Downgraded RCTs or upgraded observational studies</td>
</tr>
<tr>
<td>Low</td>
<td>Further research is very likely to have an important impact on our confidence in the estimate of effect and its likely to change the estimate</td>
<td>Well-done observational studies with control groups</td>
</tr>
<tr>
<td>Very low</td>
<td>Any estimate of effect is very uncertain</td>
<td>Others (e.g., case reports or case series)</td>
</tr>
</tbody>
</table>
The summary of the grading recommendation is summarized in table 3 and the text below summarizes the most important data for each compound.

### 1. Avocado/soybean unsaponifiables

Three RCTs have been selected for our recommendations.

The first one is a 3-month, prospective, randomized, double-blind, placebo-controlled trial evaluating the efficacy of avocado/soybean unsaponifiables (300 mg/day) in terms of NSAID use reduction [34]. Secondary efficacy criteria were the visual analogic scale pain score and the Lequesne index. After 3 months, the functional index showed a significantly greater improvement in the active drug group (-2.3 +/- 2.6) than in the placebo group (-1.0 +/- 2.6) (p < 0.01). However, pain scores over time were similar in the two groups.

The second study (n = 260), of the same duration, showed that avocado/soybean unsaponifiables (300 or 600 mg/day) significantly reduced the Lequesne index (secondary outcome) compared to placebo [35]. After 6 months of follow-up, the mean (SD) Lequesne score decreased from 9.6 +/- 2.5 to 5.5 +/- 3.6 in the avocado/soybean unsaponifiables 300 mg/day, from 9.8 +/- 2.7 to 6.5 +/- 3.5 in the avocado/soybean unsaponifiables 600 mg/day and from 9.8 +/- 2.4 to 7.8 +/- 3.4 in the placebo group (p < 0.01 between placebo and the two avocado/soybean unsaponifiables groups).

The third RCT included 164 patients with primary OA of the knee (n = 114) or hip (n = 50) with a 6-month treatment period and a 2-month post-treatment follow-up [36]. The results showed that avocado/soybean unsaponifiables (300 or 600 mg/day) significantly reduced the Lequesne index (primary outcome) and the pain score (VAS scale) compared to placebo. After 6 months of follow-up, the mean (SEM) Lequesne score decreased from 9.7 +/- 0.3 to 6.8 +/- 0.4 in the avocado/soybean unsaponifiables and from 9.4 +/- 0.3 to 8.9 +/- 0.4 in the placebo group (p < 0.001). Pain decreased from 56.1 +/- 1.6 mm to 35.3 +/- 2.3 in the avocado/soybean unsaponifiables.

In these 3 trials, avocado/soybean unsaponifiables were well tolerated.

### 2. Chondroitin sulfate

Four meta-analyses have been selected. However, two of them, showing a significant effect of chondroitin sulfate compared to placebo, were outdated since they were issued before the appearance of major recent trials [37,38]. One of them is also difficult to use since the authors evaluated chondroitin sulfate together with glucosamine sulfate [39].

In the last meta-analysis, 20 trials (3846 patients) were included [40]. The meta-analysis identified a significant beneficial effect of chondroitin sulfate on pain, with an effect size of -0.75 (-0.99 to -0.50). However, the heterogeneity between trials was high (I² = 92%). When the authors restricted the analysis to the 3 trials with large sample sizes and an intention-to-treat analysis, the effect size was -0.03 (-0.13 to 0.07; I² = 0%) and corresponded to a difference of 0.6 mm on a 10-cm visual analogic scale. However, this restricted analysis included one study with an exceptionally high placebo response rate, one study that was only published as an abstract.

At last, a meta-analysis of 12 trials showed a pooled relative risk of 0.99 (0.76 to 1.31) for any adverse event between chondroitin sulfate and placebo [40].

### 3. Diacerein

Two meta-analyses have been included.

The first one included 19 studies (search date 1985–2004) [41]. Diacerein was significantly superior to placebo to reduce pain and improve function during the active treatment phase (Glass score 1.50 [0.80–2.20]). Moreover, diacerein showed a carryover effect, persisting up to 3 months after treatment, with a significant analgesic-sparing effect during the follow-up period (Glass score 2.06

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**Table 3: Recommendations taking into account the balance of benefit (pain reduction and function improvement) and harm (adverse event)**

<table>
<thead>
<tr>
<th>Product</th>
<th>Grade of recommendation</th>
<th>Quality evidence</th>
<th>Balance benefit to harm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avocado/soybean unsaponifiables</td>
<td>Strong</td>
<td>Moderate</td>
<td>Avocado/soybean unsaponifiables advantageous</td>
</tr>
<tr>
<td>Chondroitin sulfate</td>
<td>Strong</td>
<td>Moderate</td>
<td>Chondroitin sulfate advantageous</td>
</tr>
<tr>
<td>Diacereine</td>
<td>Strong</td>
<td>Moderate</td>
<td>Diacereine advantageous</td>
</tr>
<tr>
<td>Glucosamine sulfate</td>
<td>Strong</td>
<td>Moderate</td>
<td>Glucosamine sulfate advantageous</td>
</tr>
<tr>
<td>Hyaluronic acid</td>
<td>Strong</td>
<td>Moderate</td>
<td>Hyaluronic acid advantageous</td>
</tr>
<tr>
<td>Oral calcitronin</td>
<td>Weak</td>
<td>Low</td>
<td>Calcitonin not advantageous</td>
</tr>
<tr>
<td>Risedronate</td>
<td>Strong</td>
<td>High</td>
<td>Risedronate not advantageous</td>
</tr>
<tr>
<td>Strontium ranelate</td>
<td>Weak</td>
<td>Very low</td>
<td>Strontium ranelate advantageous</td>
</tr>
</tbody>
</table>
Nevertheless, it should be pointed out that heterogeneity was not tested in this meta-analysis.

The second one included 7 studies (search date 1966–2004) with 2069 participants [42]. The results of the meta-analysis showed a small, consistent, beneficial effect of diacerein in the treatment of OA. When compared to placebo, pain on a visual analog scale (0–100 mm) showed a statistically significant difference in favour of diacerein (weighted mean differences of -0.51 [-0.96 to -0.05]); but the heterogeneity analysis result was important (p = 0.04). No significant effect of diacerein was observed on the Lequesne index for function compared to placebo, with homogeneity in all results. The most frequent adverse event was diarrhoea. 459 participants among 1083 participants that received diacerein (42%) were affected. 18% in the treatment group compared with 13% in the placebo group withdrew due to adverse events.

4. Glucosamine sulfate

Five meta-analysis and systematic reviews were included. Three groups of systematic reviews and meta-analyses emerged:

- A Cochrane Review, first published in 2001 and updated in 2005 [43], that includes every relevant glucosamine trials but studies published in 2006 and 2007. This meta-analysis remains the only one that includes all reference (NSAIDs)-controlled trials. In addition, it provides an evaluation on safety aspects.

- Meta-analyses by a group in Boston. The first was published in 2000 and is clearly outdated since it was issued before the appearance of all recent and most relevant trials [38]. However, Vlad et al. updated their meta-analysis in 2007 and this included all relevant placebo-controlled glucosamine trials [44].

- Systematic reviews and meta-analyses performed by the group in Liege. The first was published in 2003 and had the merit to be the first to include the glucosamine sulfate long-term trials [39]. However, aside not being up to date, it is also difficult to use for the assessment of glucosamine sulfate studies on the symptoms of OA, since these trials are evaluated together with those of chondroitin sulfate. Nevertheless, Reginster updated this group’s systematic review and meta-analysis in 2007 in an editorial of the meta-analysis by Vlad et al., using in his approach the pivotal trials of prescription glucosamine sulfate [44,45].

Three systematic reviews and meta-analyses have been used in the present assessment [43-45].

In the Cochrane review, the 20 analyzed RCTs found glucosamine favoured placebo with a 28% (change from baseline) improvement in pain (standardized mean difference of -0.61 [-0.95 to -0.28] and a 21% (change from baseline) improvement in function using the Lequesne index (standardized mean difference of -0.51 [-0.96 to -0.05]) [43].

Vlad et al. also reported a significant effect of glucosamine sulfate for pain improvement (effect size of 0.35 [0.14 to 0.56]) [44]. However, heterogeneity was high among the 15 RCTs included in their meta-analysis (I² = 80%).

Reginster based his meta-analysis only on 3 specific pivotal trials of glucosamine sulfate [45]. Pivotal trials are high-quality studies used by Health Authorities to assess the efficacy and safety of a prescribed medication in order to grant the marketing authorisation. The assessments of WOMAC pain and WOMAC function provide a significant beneficial effect of glucosamine sulfate compared to placebo (effect size of 0.27 [0.12 to 0.43] and 0.33 [0.17–0.48], respectively), without heterogeneity (I² = 0%).

At least, glucosamine sulfate was as safe as placebo, in terms of subjects reporting adverse reactions (RR = 0.97 [0.88 to 1.08]) [43].

5. Hyaluronic Acid

Seven systematic reviews have been included in this assessment.

Analysis of these meta-analyses showed discrepancies between systematic reviews/meta-analyses of the efficacy and safety of hyaluronic acid (HA) therapy in the treatment of osteoarthritis [46-52].

Out of the 11 RCTs included in a meta-analysis, it has been demonstrated that the 100-mm visual analog scale differences between therapy and placebo injection was 4.4 (1.1 to 7.2) at 1 week, 17.7 (7.5 to 28.0) at 5 to 7 weeks, 18.1 (6.3 to 29.9) at 8 to 12 weeks, and 4.4 (-15.3 to 24.1) at 15 to 22 weeks [51]. Another meta-analysis, including 22 RCTs, showed that patients who received the intervention treatment experienced a reduction in pain during movement: the mean difference on a 100-mm visual analog scale was -3.8 mm (-9.1 to 1.4 mm) after 2–6 weeks, -4.3 mm (-7.6 to -0.9) after 10–14 weeks and -7.1 mm (-11.8 to -2.4) after 22–30 weeks [47]. However, this effect was not considered as being clinically meaningful [47]. Another meta-analysis showed that, from the 22 studies included, the pooled effect size for hyaluronic acid was 0.32 (0.17 to 0.47) when considering pain reduction but with a significant heterogeneity among studies (P < .001) [48].

Interestingly, the reasons for inconsistency between systematic reviews have been recently searched by J. Camp-
bell et al. [57]. They identified inclusion of different controlled trials as a result of different search strategies and selection criteria, differences in the outcome measures and time points selected for extraction; and different statistical methods for data synthesis, which resulted in conflicting estimates of therapeutic effect. Anyway, the authors concluded that although the overall quality was moderate, there were net benefits (pain reduction and physical function) in favour of HA compared to placebo with low risk of harm.

6. Oral calcitonin
One RCT has been included. In this small randomized, double-blind trial, patients received either placebo (n = 18), 0.5 mg of oral salmon calcitonin (n = 17), or 1 mg of oral salmon calcitonin (n = 18) daily for 84 days [53]. No significant improvements were observed at the end of the study between patients on placebo or oral calcitonin, in the Lequesne index (secondary outcome).

7. Risedronate
Two RCTs have been included in this analysis.

The first trial included 2483 patients (placebo, 5 or 15 mg risedronate) in a 2-year study [54]. No significant effect of risedronate has been observed in the WOMAC score, compared to placebo. No increase in the number of adverse events was demonstrated for risedronate compared with placebo.

In the second study, 285 patients were randomized to once-daily risedronate (5 mg or 15 mg) or placebo, in a 1-year prospective, double-blind, placebo-controlled trial [55]. Those receiving risedronate showed no improvement of the WOMAC index, compared to placebo. Both doses of risedronate were well tolerated.

8. Strontium ranelate
One RCT has been included.

This RCT is a post-hoc analysis of two trials aiming at assessing the efficacy and safety of strontium ranelate in the treatment of postmenopausal osteoporosis [56]. The results showed that among 399 osteoporotic women with concomitant radiological spinal OA, significantly more patients in the strontium ranelate group experienced an improvement in back pain after 3 years (secondary analysis), compared with placebo (p = 0.03).

Discussion
In the light of these results, some SYSADOA have a positive risk benefit balance for patients with OA. As a matter of fact, chondroitin sulfate, diacereine, glucosamine sulfate, avocado/soybean unsaponifiables and hyaluronic acid have demonstrated pain reduction and physical function improvement with very low toxicity, with moderate to high quality evidence. The only treatment with a substantial highest level of adverse events was diacereine. The most frequent adverse effect was mild to moderate diarrhoea, which usually appeared at an early stage during treatment and resolved on continuing treatment. However, this adverse event did not result in treatment interruption in the majority of the patients.

Based on our research, there are only two treatments that are "weakly" recommended. Indeed, even if pre-clinical data and some preliminary in vivo studies have suggested that oral calcitonin and strontium ranelate could be of potential interest in OA, additional well-designed studies are needed.

It should be pointed out that some meta-analysis, even with positive results for the treatment compared to placebo, have strong evidence of heterogeneity and, with consequences, a lowest quality evidence for the treatment, following the GRADE recommendation (e.g. reduced from "high" to "moderate").

Results of the GAIT trial are also of interest and deserve special comments as this study involves glucosamine and chondroitin [58]. This National Institutes of Heath sponsored study examined placebo versus glucosamine hydrochloride (500 mg three times daily) versus chondroitin sulfate (400 mg three times daily) versus the combination of glucosamine and chondroitin versus celecoxib (200 mg/day) in a parallel, blinded 6 month multicenter study of response in knee OA. The primary efficacy variable was a 20% improvement in knee pain from baseline to 24 weeks. Overall, glucosamine and chondroitin sulfate were not significantly better than placebo in reducing knee pain by 20 percent. However, for patients with moderate-to-severe pain at baseline, the rate of response (OMERACT-OARSI criteria) was significantly higher with combined therapy than with placebo (79.2% vs. 54.3%, P = 0.002). The high placebo response (60.1%) is of unknown significance but might explain the findings of the GAIT trial. As a matter of fact, if placebo is effective in 60 percent of patients, it could be difficult for other treatments to surpass this mark. At least, this study used the glucosamine hydrochloride 500 mg three times daily compared to glucosamine sulfate 1500 once daily in the most positive trials.

It should be acknowledged, however, that the size effect in pain and physical improvement is only considered from small to moderate. At least, the duration of the RCT differs widely (3 months to 3 years), within and between treatments, making the interpretation sometime more difficult.
It should also be pointed out that some potential treatments have only been assessed in one site and that, partly because of the difference in the physiopathology between hip and knee osteoarthritis, results obtained at the level of the knee cannot be extrapolated to the hip. For example, it has been shown in one study that glucosamine sulfate appears to be ineffective in hip osteoarthritis [59].

At least, although we have used the GRADE approach to rate the quality of evidence and strength of recommendation, the need for judgment is still required. Indeed, RCTs or meta-analysis of the same product could have important methodological differences that may impact on the results. At least, even if the use of risk/benefit ratio is of great potential interest, it still needs further validation. It should also be pointed out, as a limitation of this work, that studies were not blinded and, consequently, some experts reviewed the quality of their own works.

Conclusion

In conclusion, in the benefit/risk ratio, the use of chondroitin sulfate, diacereine, glucosamine sulfate, avocado/soybean unsaponifiables and hyaluronic acid could be of potential interest for the symptomatic management of OA.

Competing interests

This article is based on the outcomes of a Working Group meeting convened by ESCEO. Some of the authors on this article have received support from, or, have given talks, attended conferences and participated in trials and advisory boards sponsored by various pharmaceutical companies. OB has received consulting fees or has been reimbursed for attending scientific meetings from Theramex, Rotta, GlaxoSmithKline, Servier, Galapados. PDD has received consulting fees, grant or has participated in paid advisory boards from Acceleron, Amgen, Eli Lilly, GSK, MSD, Novartis, Nycomed, Organon, Pfizer, Procter & Gamble, Roche, Sanofi-Aventis, Servier, Wyeth, Zelos. RR has received consulting fees, lectures fees, grant or has participated in paid advisory boards from Servier, Novartis, Amgen, GlaxoSmithKline, Roche, Nycomed, Procter & Gamble, Merck Sharp and Dohme, Lilly. CC has received consulting fees, lectures fees, grant or has participated in paid advisory boards from Servier, Proctor & Gamble/Alliance for better bone health, Merck Sharp & Dohme, Eli Lilly, GSK/Roche. JYR has received consulting fees, lectures fees, grant or has participated in paid advisory boards from Servier, Novartis, Negma, Lilly, Wyeth, Amgen, GlaxoSmithKline, Roche, Merckle, Nycomed, NPS, Merck Sharp and Dohme, Theramex, Rottapharm, IBSA, Genevriër, Teijin, Teva, Ebewee Pharma, Zodiac, Analis, Novo-Nordisk, Bristol Myers Squibb. However, the authors confirm that no pharmaceutical company has been involved with the drafting and publication of this article, financially or otherwise.

Authors’ contributions

All authors have participated in the working group meeting. OB drafted the manuscript. All authors reviewed and approved the manuscript.

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