#### SYSTEMATIC REVIEW



# Safety of Intra-articular Hyaluronic Acid Injections in Osteoarthritis: Outcomes of a Systematic Review and Meta-Analysis

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#### Abstract

**Background** Some controversy exists regarding the safety of intra-articular hyaluronic acid (IAHA) in the management of osteoarthritis (OA).

**Objective** The objective of this study was to re-assess the safety profile of IAHA in patients with OA, through a comprehensive meta-analysis of randomized, placebo-controlled trials.

**Methods** A comprehensive literature search was undertaken in the databases MEDLINE, Cochrane Central Register of Controlled Trials (CENTRAL), and Scopus. Randomized, double-blind, placebo-controlled, parallel-group trials that assessed adverse events (AEs) with IAHA in patients with OA were eligible for inclusion. Authors and/or study sponsors were contacted to obtain the full report of AEs. The primary outcomes were overall severe and serious AEs, as well as the following MedDRA System Organ Class (SOC)-related AEs: gastrointestinal, cardiac, vascular, respiratory, nervous system, skin and subcutaneous tissue disorders, musculoskeletal, renal and urinary disorders, infections and infestations, and hypersensitivity reaction.

**Results** Database searches initially identified 1481 records. After exclusions according to the selection criteria, 22 studies were included in the qualitative synthesis, and nine studies having adequate data were ultimately included in the metaanalysis. From the studies excluded according to the pre-specified selection criteria, 21 with other pharmacological OA treatments permitted during the trials were a posteriori included in a parallel qualitative synthesis, from which eight studies with adequate data were finally included in a parallel meta-analysis. Since this meta-analysis was designed to assess safety, the exclusion criterion on concomitant anti-OA medication was crucial. However, due to the high number of studies that allowed mainly concomitant oral non-steroidal anti-inflammatory drugs (NSAIDs), we decided to include them in a post hoc parallel analysis in order to compare the results from the two analyses. No statistically significant difference in odds was found between IAHA and placebo for all types of SOC-related disorders, except for infections and infestations, for which significantly lower odds were found with IAHA compared with placebo, both overall (odds ratio [OR]=0.61, 95% confidence interval [CI] 0.40–0.93;  $I^2$ =0%) and in studies without concomitant anti-OA medication (OR=0.49, 95% CI 0.27–0.89). There were significant increased odds of reporting serious AEs with IAHA compared with placebo, both overall (OR=1.78, 95% CI 1.10–2.89), but not in studies without concomitant anti-OA medication (OR=1.78, 95% CI 1.10–2.89), but not in studies without concomitant anti-OA medication (OR=1.78, 95% CI 1.10–2.89), but not in studies without concomitant anti-OA medication (OR=1.78, 95% CI 1.10–2.89), but not in studies without concomitant anti-OA medication (OR=1.78, 95% CI 1.20–3.47).

**Conclusions** Using the available data on studies without any concomitant anti-OA medication permitted during clinical trials, IAHA seems not to be associated with any safety issue in the management of OA. However, this evidence was associated with only a "low" to "moderate" certainty. A possible association with increased risk of serious AEs, particularly when used with concomitant OA medications, requires further investigation.

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Extended author information available on the last page of the article

#### **Key Points**

Our meta-analysis of randomized controlled trials (RCTs) without any concomitant pharmacological osteoarthritis (OA) treatments permitted during the trials did not identify any safety issue associated with intraarticular hyaluronic acid (IAHA); however, the certainty of this new evidence was graded between "low" and "moderate".

A shortcoming in the reporting of harms-related data in manuscripts communicating outcomes of RCTs with IAHA in OA was the reason for this uncertainty, which does not allow for a definitive conclusion regarding the safety of IAHA.

Additional studies are required to further investigate the safety profile of IAHA, particularly any association with serious adverse events and long-term safety; moreover, authors of studies on IAHA are encouraged to report in a transparent way all harms data collected from RCTs in the future.

### 1 Introduction

Osteoarthritis (OA) is a chronic disorder, affecting joints such as hand, knee and hip, that causes considerable pain, increasing disability, and progressive cartilage degeneration [1]. OA occurs frequently in adults aged over 50 years and is a major cause of disability worldwide [2, 3]. The incidence of OA is rising due to the aging population and the increase in obesity [1]. OA has a complex pathophysiology that is incompletely understood. There is no established diseasemodifying therapy for OA as yet, and hence the treatment of OA relies on a combination of pharmacological and nonpharmacological therapies that can manage the symptoms of OA, primarily pain and loss of function [4]. In practice, paracetamol, non-steroidal anti-inflammatory drugs (NSAIDs) and opioids are widely prescribed to relieve pain and improve joint function, and yet have significant toxicity [5-8]. Analgesics and NSAIDs are particularly poorly tolerated by OA patients [9, 10], who are frequently of advanced age, with comorbidities and are receiving polypharmacy. Intra-articular (IA) injections of corticosteroids, such as triamcinolone hexacetonide and methylprednisolone acetate, are also often prescribed. However, systemic absorption occurs following IA corticosteroid injection, which can lead to systemic adverse events (AEs), and precautions should be observed in patients with concomitant diseases, such as hypertension and diabetes mellitus [11–13].

Intra-articular hyaluronic acid (IAHA) injection presents an alternative local treatment option providing symptomatic benefit without the systemic AEs associated with IA corticosteroids. Numerous RCTs and meta-analyses have sought to assess the efficacy and safety of IAHA, with mixed results and conclusions [14, 15]. IAHA is demonstrated to have a positive effect on pain and joint function [16]. A network meta-analysis comparing the effectiveness of pharmacological interventions for knee OA found IAHA to be the most efficacious treatment, with an effect size (ES) of 0.63 on pain (95% credible interval [CrI] 0.39-0.88). IA placebo itself had an ES of 0.29 (95% CrI 0.04-0.54); nonetheless a statistically significant effect for IAHA was found at 3 months (ES = 0.34, 95% CrI 0.26–0.42) [17]. IAHA is also demonstrated to have a longer lasting effect on pain and function compared with IA corticosteroids, lasting up to 6 months [18, 19].

There is also mounting data showing that multiple courses of IAHA can impact long-term outcomes, including reduction in concomitant analgesia use, and delay in the need for total knee replacement surgery [20–22]. However, despite evidence attesting to the efficacy of IAHA injections, particularly for knee OA, and the widespread use of IAHA in clinical practice, controversy still persists regarding the relative risk:benefit of IAHA, largely due to mixed reports on the safety of IAHA. Consequently, there is a lack of agreement among national and international guidelines regarding the use of IAHA for the medical management of symptomatic knee OA [4, 23–29].

Eight meta-analyses of RCTs comparing IAHA to IA placebo have evaluated the safety of IAHA [16, 17, 30-35]. A Cochrane review of 76 RCTs was unable to conclude a definitive comment on the safety of the hyaluronic acid (HA) class of products due to sample-size restrictions; however, no major safety issues were detected, and in some analyses, IAHA demonstrated similar efficacy to systemic forms of active intervention, with more local reactions but fewer systemic AEs [16]. Recent meta-analyses on the safety of different IAHA products have found that HA is generally well-tolerated, with a low incidence of AEs and without risk for serious adverse events (SAEs) [35-37]. However, a meta-analysis published in 2012 raised concerns regarding the safety of IAHA, finding an increased risk of SAEs associated with IAHA compared with sham or non-intervention control (relative risk = 1.41, 95% confidence interval [CI] 1.02–1.97) [33]. Notably, almost all of those previous meta-analyses which have assessed the safety of IAHA used only published data, and it is well known that safety data are under-reported in manuscripts. Additionally, the concomitant use of oral NSAIDs during some clinical trials was not taken into account in the previous analyses. The objective of this study was therefore to reassess the safety

of HA injections in the management of OA in a systematic review and meta-analysis of randomized, placebo-controlled trials (RCTs). In order to better estimate the safety profile of IAHA in OA, authors of the manuscripts and/or sponsors of studies were contacted to get the full report of AEs.

# 2 Methods

The protocol of this systematic review and meta-analysis has been previously registered in the PROSPERO database (registration number: CRD42017071906). The systematic review was performed in accordance with the recommendations in the *Cochrane Handbook for Systematic Reviews of Interventions* [38]. The findings were reported according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines [39]. All the review process (study selection and risk of bias assessment) was undertaken using Covidence, the Cochrane platform for systematic reviews.

#### 2.1 Eligibility Criteria

Randomized, double-blind, placebo-controlled, parallelgroup trials which have assessed AEs associated with IAHA in patients with OA were eligible for inclusion in this metaanalysis. The following studies were excluded: cross-over studies, reviews or meta-analyses, letters, comments or editorials. Studies that allowed concomitant anti-OA medications during the trial (other than rescue medication such as paracetamol or aspirin) were also excluded for the main meta-analysis, but were kept and used for a parallel analysis.

#### 2.2 Data Sources and Search Strategies

A comprehensive literature search was undertaken in the databases MEDLINE (via Ovid), Cochrane Central Register of Controlled Trials (Ovid CENTRAL) and Scopus. Each database was searched from inception up until 19 June 2017. We searched for RCTs of IAHA in OA, using a combination of study design-, treatment-, and disease-specific key words and/or Medical Subject Heading (MeSH) terms. While AEs were the outcomes of interest for this study, we decided to avoid the outcome-specific key words in the search strategies because of the possibility that a study on the efficacy of a drug may have not mentioned terms related to AEs in its title, abstract or in the keywords sections. The search was limited to English and French publications and to human subjects. Detailed search strategies for MEDLINE/CEN-TRAL and Scopus databases are reported in the Electronic Supplementary Material (ESM1).

Two clinical trial registries, ClinicalTrials.gov (http:// clinicaltrials.gov/) and the World Health Organization's International Clinical Trials Registry Platform Search portal (http://apps.who.int/trialsearch/), were also checked for trial results that would not have been published. Finally, very recent meta-analyses were also screened for any additional relevant studies. For all studies that met the selection criteria, authors of the manuscripts and/or sponsors of studies were automatically contacted to get the full report of AEs, as far as there was any way to contact them (email, fax, telephone number or co-author email in another article).

We set up search alerts in the bibliographic databases for any new relevant RCTs that were published from 19 June 2017 until 30 September 2018.

#### 2.3 Study Selection and Data Extraction

Two members of the review team (GH and XR) independently evaluated each title and abstract to exclude only obvious irrelevant studies, according to the predefined eligibility criteria. At this step, the criteria related to adverse effects were not considered for selection, as studies focusing on the efficacy of a treatment may not report data about adverse effects in the abstract; this means that all trials mentioning only the efficacy information were retrieved at this step. After this first step, the two investigators independently reviewed the full texts of the articles not excluded during the initial screening stage to determine whether the studies met all selection criteria. Those which did not meet these criteria were definitely excluded. All differences of opinion regarding the selection of articles were resolved through discussion and consensus between the two investigators (GH and XR); any persistent disagreement was solved with the intervention of another member of the review team (AG or VR). A flowchart with the number of included studies at each step was established, including the reasons for excluding studies during the full-text reading process.

The full texts of the selected studies were screened for extraction of relevant data, using a standard data extraction form. Outcome results data were independently extracted by two investigators of the review team (GH and XR). For each study, the following data were extracted: characteristics of the manuscript, characteristics of the trial, objective and design of the study, characteristics of the patients, characteristics of the disease, characteristics of the treatments, AEs (outcomes) reported during the trial, and the main conclusion of the study. The raw data (number of events in each arm of the study) were extracted for each outcome. The number of patients who experienced at least once any body system related AE (e.g. nervous system, gastrointestinal system) as well as specific AEs within each body system (e.g. headache, abdominal pain) were extracted. The total number of patients who experienced at least once any AE during the trial and the total number of patients who withdrew from the trial due to AEs were also extracted. Intention-to-treat data were only used when reported or supplied by the study authors or sponsor.

# 2.4 Assessment of Risk of Bias in Included Studies

Two members of the review team (GH and XR) independently assessed the risk of bias in each study, using the Cochrane Collaboration's tool for risk of bias assessment [38]. The following characteristics were evaluated:

- Random sequence generation: we assessed whether the allocation sequence was adequately generated.
- Allocation concealment: we assessed the method used to conceal the allocation sequence, evaluating whether the intervention allocation could have been foreseen in advance.
- Blinding of participants and personnel: we assessed the method used to blind study participants and personnel from knowledge of which intervention a participant received and whether the intended blinding was effective.
- Blinding of outcome assessment: we assessed the method used to blind outcome assessors from knowledge of which intervention a participant received and whether the intended blinding was effective.
- Incomplete outcome data: we assessed whether participants exclusions, attrition and incomplete outcome data were adequately addressed in the paper.
- Selective outcomes reporting: we checked whether there was evidence of selective reporting of AEs.

Each of these items was either categorized as "low risk of bias", "high risk of bias", or "unclear risk of bias". "Low risk of bias" or "high risk of bias" was attributed for an item when there was sufficient information in the manuscript to judge the risk of bias as "low" or "high"; otherwise, "unclear risk of bias" was attributed to the item. Disagreements were solved by discussion between the two reviewers (GH and XR) during a consensus meeting and involved, when necessary, another member of the review team for final decision (AG or VR).

# 2.5 Outcomes of Interest

The main System Organ Classes (SOCs) that are likely to be affected by the use of IAHA in the treatment of OA were explored in this meta-analysis. The primary outcomes of interest were MedDRA (Medical Dictionary for Regulatory Activities) SOC-related AEs: gastrointestinal disorders, cardiac disorders, vascular disorders, respiratory, thoracic and mediastinal disorders, nervous system disorders, skin and subcutaneous tissue disorders, musculoskeletal and connective tissue disorders, renal and urinary disorders, infections and infestations, as well as hypersensitivity reaction, and overall severe AEs and SAEs. Secondary outcomes were withdrawals due to AEs (i.e. the number of participants who stopped the treatment due to an AE) and total number of AEs (i.e. the number of patients who experienced any AE at least once).

# 2.6 Data Analysis

Analyses were performed using STATA 14.2 software. We described harms associated with the treatment as odds ratio (OR) with 95% CI. We computed an overall ES for each primary or secondary outcome (AE). Anticipating substantial variability among trial results (i.e. the inter-study variability), we assumed heterogeneity in the occurrence of the AEs; thus, we planned to use random-effects models for the meta-analyses. We estimated the overall effects and heterogeneity using the DerSimonian and Laird random-effects model [40]. As this method provides a biased estimate of the between-study variance with sparse events [41, 42], we also performed the meta-analyses using the restricted maximum likelihood (REML) method [43]. Indeed, we planned in the protocol to use specific methods for rare events analysis, in case it would be necessary. However, we reported only the results from the DerSimonian and Laird random-effects model, as we found no difference in the effects computed by the two methods. We preferred reporting the results from the DerSimonian and Laird method (which uses a correction factor), because it allows for displaying studies with null event on the forest plot, even if those with null event in both the intervention and control groups are excluded from the overall ES computation. On the contrary, with the REML method, these studies are not displayed on the forest plot. Additionally, the STATA command which performs the meta-analysis based on the REML method (metaan) does not have any option for displaying subgroups on the same graphic, in contrary to the DerSimonian and Laird method command (metan), which has this option ("by").

We tested heterogeneity using the Cochran's Q test. As we are performing a random-effects meta-analysis, we used the Tau-squared (Tau<sup>2</sup>) estimate as the measure of the between-study variance. The *I*-squared ( $I^2$ ) statistic was used to quantify heterogeneity, measuring the percentage of total variation across studies due to heterogeneity [44]. In the case of substantial heterogeneity ( $I^2 > 50\%$ ) [45], we pre-specified to undertake subgroup analyses, stratifying the analyses according to the following: participants' age in the intervention group, duration of OA complaint, type

of joint treated (knee, hip), number of joints treated, origin of the HA used (avian, microbial), molecular weight of HA (high weight versus low weight), molecular structure of HA (cross-linked versus non-cross-linked), HA manufacturer, dosing regimen, number of cycles, number of injection per cycle (single versus repeated), follow-up duration, type of sham intervention used (saline versus other), use of anaesthetic before injection, ultrasound guidance for injection, industry involvement (sponsored versus non-sponsored), risk of bias in the studies (e.g. studies with low risk of bias versus all other studies).

We assessed evidence for publication bias by visual inspection of funnel plots and using the Harbord's test for funnel plot asymmetry [46], which is more suitable for dichotomous outcomes with ESs measured as OR [47] than the classical Egger's test [48]. The quality of each evidence was assessed using the GRADE approach [49], and a summary of findings table was prepared using the GRADEpro online software [50].

#### 2.7 Additional Analysis

We performed additional post hoc meta-analyses, in parallel with the main meta-analysis including the studies responding to our pre-defined eligibility criteria. Indeed, studies allowing concomitant anti-OA medications, as excluded based on our eligibility criteria, as well as all studies with or without concomitant anti-OA medications, were considered separately in parallel to the primary meta-analysis. These parallel analyses were done based on the same principles announced in the data analysis section of this manuscript for the main meta-analysis. However, instead of depicting the results of the parallel analyses in separate forest plots, we preferred showing all the analyses for each outcome on the same figure, to allow for an easy comparison. Therefore, considering the rationale of this safety meta-analysis (the exclusion of studies with other anti-OA medication allowed), the parallel analyses on one single forest plot are not to be considered as subgroup analyses, as for a classical meta-analysis.

#### **3 Results**

#### 3.1 Initial Study Selection and Characteristics

Database searches initially identified 1481 records, from which, after exclusions, 88 articles were assessed for eligibility. Of these, 67 studies were excluded for various reasons (Fig. 1). Twenty-two papers were included in the qualitative synthesis, according to our pre-specified selection criteria, and only nine studies having adequate data were ultimately included in the meta-analysis [51–72]. These studies responding to our selection criteria were without any concomitant pharmacological OA therapy, in accordance with the review protocol. Indeed, since this is a meta-analysis on the safety of IAHA, we could not include trials that had allowed another pharmacological OA treatment as concomitant medication (other than rescue medication such as paracetamol or aspirin).

Table 1 presents the characteristics of the studies included through the systematic review process (those included in the quantitative synthesis—meta-analysis—are highlighted). Almost all these studies involved patients with knee OA; only three were on patients with other joint OA (ankle OA). Most of the trials had follow-up durations varying between 25 and 29 weeks; only one trial had a follow-up duration of at least 52 weeks. Single or repeated injections of HA during a single cycle were reported, respectively, by four and 17 studies; one study was a multiple-arm study with single and repeated injections (one, three or five injections). Avian derivative or microbial derivative HA was used, but in the majority of the articles, the origin of the compound was not specified.

Of the 22 articles selected for inclusion, only seven had their published data usable for the meta-analysis; thus, the risk of selective outcome reporting was found to be "high" in more than 75% of the retrieved studies. Full safety data were received for only two studies (Table 1). Figures 2a and 3a include a summary of the risk of bias assessed for each study included in the qualitative synthesis and the risk of bias items presented as percentages across all these studies.

#### 3.2 Post Hoc Study Selection and Characteristics

From the 67 studies previously excluded according to the protocol, 21 with other pharmacological OA treatments permitted were "a posteriori" included in a parallel qualitative synthesis (Fig. 1), from which eight studies with adequate data were ultimately included in a parallel meta-analysis [73–93]. In most of these studies, oral NSAIDs were permitted as rescue or concomitant medication; in a few others, opiates and steroids were allowed.

This "post hoc" decision to consider these studies with other pharmacological OA treatments in a parallel analysis was made because we were surprised by their number when compared to the number of studies without any concomitant pharmacological OA treatment allowed. We sought by so doing to compare the results from these two groups of studies, knowing that our main conclusions regarding the safety profile of IAHA will primarily be based on the results of the analyses using the studies without any concomitant pharmacological OA treatment (those responding to our prespecified selection criteria).



Fig. 1 Flow chart of the study. OA osteoarthritis

Table 2 presents the characteristics of the studies included in the parallel qualitative synthesis (those included in the meta-analysis are highlighted). The majority of the studies retrieved for this parallel qualitative synthesis included patients with knee OA, as for the studies without any concomitant anti-OA medication; three studies included patients with hip OA, one studied patients with hand OA, and two others each included patients with OA of the glenohumeral joint and the first metatarsophalangeal joint. For most of the studies, the follow-up durations were  $\leq 26$  weeks; three studies had follow-up durations of 52 weeks and one of 174 weeks. From the 21 studies included in the parallel qualitative synthesis (i.e. with concomitant anti-OA medication), only eight, with data partially reported, were included in the parallel quantitative synthesis (meta-analysis); for the others, the published data were not usable for the meta-analysis and no adequate data were obtained from the study authors or sponsors. A "high" risk of selective outcome reporting bias was found in the majority of the studies. Figures 2b and 3b include a summary of the risk of bias assessed for each study included in the parallel qualitative synthesis and the risk of bias items presented as percentages across all these studies.

Table 1         Chara           included in the	cteristics c meta-anal	of the studies included fi ysis are highlighted in b	rom the systematic 1 oold type)	review process,	according to the pre-specified sel	ection criteria (w	ithout concomitant OA	treatment; those	studies ultimately
Study	Location of OA	Treated groups/age of participants (mean ± SD or median [P25- P75])	Origin of HA	Molecular weight	Dose Number of cycles/number of injections per cycle	Follow-up dura- tion (weeks)	Data provided in the article (type of AE/% of patients consid- ered)	Published data usable for M-A? (yes/no)	Full data provided by the author/ sponsor? (source of information)
Altman 1998 [51]	Knee	IA HA: 62 ± 10 IA SA: 65 ± 10	Avian sodium hyaluronate	500–730 kDa	2 mL (20 mg) HA 1/5 in saline vehicle	26 weeks	Most commonly occur- ring AEs (> 5%) on all randomized patients	Yes	No (Fidia Pharma)
Altman 2004 [52]	Knee	IA HA: Mean: 62.9 Range : 41–85 IA SA: Mean: 63.3 Range: 35–85	Streptococci [Non- animal stabilized hyaluronic acid (NASHA)]	NA	3 mL injection : 1/1 60 mg HA in buffered sodium chloride, 0.9%	26 weeks	Summary according to the relationship of AEs to the treatment. Data for TRAEs (the most common) and serious AEs	Yes	No (Author)
Arden 2014 [53]	Knee	IA HA: 64.5 (29–84) IA SA: 60.9 (30–86)	NA	NA	60 mg, single IA 1/1 injection	6 weeks	TRAEs/Summary	No	No (Author)
Baltzer 2009 [54]	Knee	IA HA: 57.4 ± 12.0 IA SA: 60.3 ± 10.7	Ч.	1.4 x 10 <sup>6</sup> Da	2 mL of HA (1% 1/3 solution of HA)	29 weeks	Results mainly pre- sented as percent- ages and numbers of patients with 'mild', 'moderate', 'severe' AEs. Frequencies reported for some SOCs	Yes	Author con- tacted with no response
Brandt 2001 [55]	Knee	IA HA: 65 ± 8.4 IA SA: 67 ± 8.4	Avian (Rooster combs)	1.0–2.9 million Da	2 mL (15 mg/mL) 1/3	27 weeks	AEs reported by $\geq 5$ % of the patients (by Body System)	Yes	Author con- tacted with no response
Carrabba 1995 [56]	Knee	IA HA 1 injection: 61.3 $\pm$ 6.8 IA HA 3 injections: 60.0 $\pm$ 7.1 IA HA 5 injections: 60.6 $\pm$ 7.9 IA Placebo: 60.0 $\pm$ 7.0	Avian (Rooster combs)	500–730 kDa	20 mg/2 mL, 1, 1/1, 3 or 5 3 or 5 injections	26 weeks	Summary; focus on local AEs.	No	No contact infor- mation found
Cohen 2008 [57]	Ankle	IA HA: 56.2 ± 15.1 IA SA: 43.4 ± 14.9	NA	500–730 kDa	NA (2 mL HYL) 1/5	26 weeks	Summary/Not detailed	No	Author con- tacted with no response
DeCaria 2012 [58]	Knee	IA HA: 71.93 ± 6.83 IA inert HA (Placebo): 72.93 ± 5.48	NA	730 kDa	2ml of 20 mg/ 1/3 ml HA	26 weeks	Summary/Not detailed.	No	Author con- tacted with no response

Table 1 (conti	nued)									
Study	Location of OA	Treated groups/age of participants (mean ± SD or median [P25– P75])	Origin of HA	Molecular weight	Dose	Number of cycles/number of injections per cycle	Follow-up dura- tion (weeks)	Data provided in the article (type of AE/% of patients consid- ered)	Published data usable for M-A? (yes/no)	Full data provided by the author/ sponsor? (source of information)
DeGroot 2012 [59]	Ankle	IA HA: 54.1 ± 14.5 IA SA: 61.9 ± 14.1	Avian derivate (derived from rooster combs)	620,000 to 1,170,000 Da	Each 2.5 mL of Supartz contains 25 mg of sodium hyaluronate	1/1	12 weeks	Summary/Not detailed	No	Author con- tacted with no response
Diracoglu 2009 [60]	Knee	IA HA: 59.4 ± 9.9 IA SA: 56.2 ± 7.2	Ч Ч	Ϋ́Υ	۷X	1/3	NA (Short-term study. 'Injec- tions were repeated in both groups three times after every one-week')	Summary/Not detailed	Ŷ	Author con- tacted with no response
Dixon 1988 [61]	Knee	IA HA and IA SA: Mean: 68.5 Range: 43–85	NA	NA	20 mg sodium hyaluronate (2 mL)	1/11	48 weeks	Treatment related or possibly related AEs reported.	Yes	No (Fidia Pharma)
Gormeli 2017 [62]	Knee	IA HA: 53.5 ± 14 IA SA: 52.8 ± 12.8	NA	NA ('A high molecular weight HA')	2 mL (30 mg/ 2 mL HA)	1/3	26 weeks	AEs assessed, but results not reported	No	Author con- tacted with no response
Hangody 2017 [63]	Knee	IA HA: 59.2 ± 8.6 IA SA: 58.0 ± 9.0	NA	NA	4 mL, 88 mg HA	1/1	26 weeks	Summary: numbers of AEs reported, not frequencies	Ŷ	Yes, data sent by the author as "Number of AEs - By Severity" and not frequencies; not adequate for M-A
Huang 2011 [64]	Knee	IA HA: 65.9 ± 8.1 IA SA: 64.2 ± 8.4	Avian (naturally derived from rooster combs)	500–730 kDa	20 mg HA /2 mL	1/5	25 weeks	Summary/Not detailed. Number of SAEs reported, not frequen- cies	No	No (Fidia Pharma)
Jorgensen 2010 [65]	Knee	IA HA: 62.6 ± 11.4 IA SA: 61.4 ± 11.1	NA	NA	2 ml Hyalgan (10 mg/ml)	1/5	13 weeks to a maximum of 52 weeks after the 1st injection	Summary/Not detailed. Per group numbers of AEs reported	°N	Yes (Author)

Table 1 (conti	inued)									
Study	Location of OA	Treated groups/age of participants (mean $\pm$ SD or median [P25– P75])	Origin of HA	Molecular weight	Dose	Number of cycles/number of injections per cycle	Follow-up dura- tion (weeks)	Data provided in the article (type of AE/% of patients consid- ered)	Published data usable for M-A? (yes/no)	Full data provided by the author/ sponsor? (source of information)
Karlsson 2002 [66]	Knee	IA HA (Artzal): 72 $\pm 7$ IA HA (Synvisc): 70 $\pm 7$ IA SA: 71 $\pm 6$	NA	Artzal : ~10 <sup>6</sup> Da Synvisc : ~7 x 10 <sup>6</sup> Da	Artzal: 2.5 ml (1% HA); Synvisc : 2.0 ml (0.8% HA)	1/3	52 weeks	Summary/Not detailed. Number of AEs reported, not the frequencies of specific AEs	No	No (Author)
Kotevoglu 2006 [67]	Knee	IA HA (Orthovisc) : 58.6 ± 8.0 IA HA (Synvisc) : 59.7 ± 8.0 IA SA : 60.1±5.4	Ч И	NA (Orthovise: Low molecular weight Synvise: High molecular weight)	A VA	1/3	26 weeks	Summary/Not detailed	°Z	Author con- tacted with no response
Kul-Panza 2010 [68]	Knee	IA HA: 59.5 ± 8.8 IA SA: 62.8 ± 7.8	Ч И	1.5 million Da (average)	2 mL 1.5% (15 mg/mL) of intra articular HA	<ul><li>1/9 (3 injections</li><li>a week, for 3 consecutive weeks).</li></ul>	14 weeks	Summary/Not detailed	No	Clarifications from the author not sufficient for inclusion of this study in the M-A
Petrella 2002 [69]	Knee	IA HA (+ Placebo tablet): $67.3 \pm 8.9$ IA SA (+ Placebo tablet): $62.6 \pm 9.5$	NA (Suplasyn)	NA	2ml, 10mg/ml	1/3	12 weeks	Summary/Not detailed	No	Author con- tacted with no response
Puhl 1993 [70]	Knee	IA HA: 62.1 (41–75) IA SA: 60.8 (40–74)	Avian (Rooster combs)	6.0–12.0 × 10 <sup>5</sup> Da	25 mg of sodium hyaluronate/ 2.5 ml	1/5	18 weeks	All AEs seem to have been reported (treatment-related and others)	Yes	No contact infor- mation found
Salk 2006 [71]	Ankle	IA HA: 57.8 ± 14.7 IA SA: 60.0 ± 13.9	NA	500–730 kDa	1 mL of hyalu- ronic acid (10 mg/mL)	1/5	26 weeks	Detailed report of AEs: usable for analysis	Yes	Author con- tacted with no response
Van der Weegen 2015 [72]	Knee	IA HA: 58.7 ± 9.6 IA SA: 60.1 ± 10.1	Produced from the bacterium <i>Strep-</i> <i>tococcus equi</i> by <i>i</i> process of continu- ous fermentation.	2.2 M Da	1.5 % HA (30 mg/2 ml)	1/3	29 weeks	Summary/Not detailed	No	Yes (Author)
Where publist sor	ied data we	re adequate for inclusio	n in the M-A and a	full safety repor	t was also provide	d by the author/s	sponsor, we pref	erentially used the full	data obtained fro	n the author/spon

Meta-Analysis of IAHA Safety in OA

S109

AE adverse event, HA hyaluronic acid, IA intra-articular, IAHA intra-articular hyaluronic acid, IASA intra-articular saline, M-A meta-analysis, NA not available (i.e. information not provided in the manuscript), OA osteoarthritis, SAE serious adverse event, SOC System Organ Class, TRAEs treatment-related adverse events

Fig. 2 a Risk of bias summary in studies without concomitant pharmacological OA treatment (studies meeting the pre-specified selection criteria): review authors' judgements about each risk of bias item for each study included in the primary qualitative synthesis. b Risk of bias summary in studies with concomitant pharmacological OA treatment (studies included in the parallel qualitative synthesis): review authors' judgements about each risk of bias item for each study included in the parallel qualitative synthesis. OA osteoarthritis



#### 3.3 Primary Outcomes

Using the available data, IAHA was found to be associated with significantly lower odds of infections and infestations, overall (OR = 0.61, 95% CI 0.40–0.93;  $l^2 = 0\%$ ) and in studies without any concomitant anti-OA medication allowed (OR = 0.49, 95% CI 0.27–0.89;  $l^2 = 0\%$ ) (Fig. 4). A reduced

odds of infections was also found with studies that allowed concomitant OA treatment, but this did not reach statistical significance (OR = 0.76, 95% CI 0.42-1.38). For this outcome, two main studies accounted for more than 96% of the weight in the overall analysis. Influenza, urinary tract infection and pneumonia were the most reported specific events in the placebo group, and these specific events were

Fig. 3 a Risk of bias graph for studies without concomitant pharmacological OA treatment: review authors' judgements about each risk of bias item presented as percentages across all studies included in the primary qualitative synthesis. b Risk of bias graph for studies with concomitant pharmacological OA treatment: review authors' judgements about each risk of bias item presented as percentages across all studies included in the parallel qualitative synthesis. OA osteoarthritis



reported by only one study, for which the authors shared the full safety report with us 65].

No statistically significant difference was found between IAHA treatment and placebo for all other types of disorders, including gastrointestinal, cardiac, vascular, respiratory, thoracic and mediastinal, nervous system, skin and subcutaneous tissue, musculoskeletal and connective tissue, renal and urinary system disorders, and hypersensitivity reaction (see the Electronic Supplementary Material, ESM2).

There were significant increased odds of reporting SAEs in the IAHA group, both overall (OR = 1.78, 95% CI 1.21–2.63;  $I^2 = 0\%$ ) and in studies that allowed concomitant OA treatment (OR = 1.78, 95% CI 1.10–2.89;  $I^2 = 0\%$ ) (Fig. 5). An increased odds of SAEs was also found in studies without concomitant anti-OA treatment, but this did not reach statistical significance (OR = 1.78, 95% CI 0.92–3.47;  $I^2 = 0\%$ ). No statistically significant difference was found between IAHA treatment and placebo for severe AEs.

#### 3.4 Secondary Outcomes

Overall, there were no more total AEs reported with IAHA versus placebo (OR = 1.09, 95% CI 0.90–1.31;  $I^2 = 17.7\%$ ) and specifically without concomitant OA treatment allowed (OR = 1.19, 95% CI 0.92–1.54;  $I^2 = 0\%$ ) and with concomitant OA treatment (OR = 1.04, 95% CI 0.77–1.40;  $I^2 = 0\%$ ) (Fig. 6).

Overall, there were significant increased odds of withdrawals due to AEs with IAHA (OR = 1.62, 95% CI 1.04–2.51;  $I^2 = 10.0\%$ ) and increased but not statistically significant odds of withdrawals due to AEs in studies of IAHA without concomitant anti-OA treatment (OR = 1.80, 95% CI 0.99–3.26;  $I^2 = 23.7\%$ ) (Fig. 7).

#### 3.5 Assessment of Publication Bias

Funnel plot asymmetry was visually investigated for each of the primary or secondary outcomes assessed for IAHA compared with placebo. The Harbord's test was also performed, when sufficient data were available. Whatever the outcome considered, visual inspection of funnel plots and the Harbord's test for funnel plot asymmetry (when possible) showed that there was no evidence of publication bias. The funnel plot for "total AEs" is depicted in Fig. 8; all the other funnel plots are provided as Electronic Supplementary Material (ESM3).

#### 3.6 GRADE Assessment of Findings

We assessed the certainty of evidence for each primary or secondary outcome for IAHA compared with placebo, using the GRADE approach [49]. Our findings were associated with "low" to "moderate" certainty of evidence, due to serious risk of bias issues (reporting bias and/or attrition bias) across the included studies. Additionally, we found large imprecisions with some overall effect estimates because of a low number of events (null events were reported in many of the included studies and for most of the outcomes). Table 3 summarizes

Table 2 Char highlighted ir	racteristics o	f studies included	l in the parallel q	ualitative synth	lesis (trials with con	ncomitant OA t	reatment allo	wed; those studies	ultimately inclu	ided in the par	allel meta-analysis are
Study	Location o OA	of Treated groups/Age of participants (mean ± SD or median [P25- P75])	Origin of HA	Molecular weight	Dose	Number of cycles/number of injections per cycle	Follow-up duration (weeks)	Concomitant OA treatment (medi- cation) allowed	Data provided in the article (type of AE/% of patients considered)	Published data usable for M-A? (yes/no)	Full data provided by the author/sponsor? (source of informa- tion)
Altman 2009 [73]	Knee	IA HA: 62,5 ± 11 IA SA: 60,8 ± 10	Derived from nonpyogenic Strepto- coccus zooepidemi- cus (biologic fermenta- tion)	2.4–3.6 Mil- lion Da	20mg/2ml of 1% solution hyalu- ronate	1/3	26 weeks	Non-prescription nutraceuticals (e.g., glucosa- mine, chon- droitin), topical analgesics, and nasal or inhaled corticosteroids	Report mainly focused on Musculoskel- etal and Con- nective Tissue AEs. Any, Withdrawals, Severe and serious AEs also reported. No detail for others	Yes	Author contacted with no response
Atchia 2011 [74]	Hip	IA HA: 69 ± 9 IA SA: 70 ± 10	NA (Non- animal sta- bilized HA, Durolane)	NA (a high molecular weight hya- luronan)	3ml/60mg	1/1	8 weeks	There were no restrictions regarding medi- cation use	Summary/Not detailed	No	Yes, data sent by the author as number of AEs, and not frequencies; not adequate for M-A.
Chevalier 2010 [75]	Knee	IA HA: 63.6 ± 9.64 9.64 9.17 9.17	NA [Hylan G-F 20 (Syn- visc)]	6000 kDa (Average)	NA (6 ml intra- articular injec- tion)	1/1	26 weeks	Analgesics/ NSAIDs (with a half-life of 5 h or less for indications other than osteoarthri- tis pain)	Report focused on target knee AEs	No	No (Author)
Day 2004 [76]	Knee	Mean (Range) IA HA: 62 (39-79) IA Placebo vehicle: 62 (33-75)	Avian (extracted from rooster combs)	6.2 × 10 <sup>5</sup> to 11.7 × 10 <sup>5</sup> Da	25 mg of sodium HA in 2.5 ml of phosphate buffered saline	1/5	17 weeks	In the results sec- tion: Impor- tant Codeine compounds and NSAID use in violation of the protocol, but patients not excluded	Summary/Not detailed	°Z	No (Author)

an of Treated     Origin of HA     Molecular     Dose     Number of groups/Age of groups/Age of median [P25-       P75])     Barticipants     weight     Dose     Number of cycles/number of injoctions       P75])     P75]     Avian     500-730 kDa     Hyalectin 20 mg     1/4       P75])     JA Vehicle     from cock's     500-730 kDa     Hyalectin 20 mg     1/4       P75])     JA Vehicle     from cock's     500-730 kDa     Hyalectin 20 mg     1/4       P75]     Avian     500-730 kDa     Hyalectin 20 mg     1/4       P75     (10 mg/ml)     1/4     1/5       P75     Data for serer     Avian     500,000 Da     20 mg Hyalgan in 1/5       P11     TAHA: 72.1     from roster     -750,000 Da     2 ml of sterile       IA Vehicle     67.0 ± 1.7     from roster     -750,000 Da     2 ml of sterile       IA Vehicle     67.0 ± 1.7     from roster     -750,000 Da     2 ml of sterile       IA Vehicle     67.0 ± 1.7     from roster     -750,000 Da     2 ml of sterile       IA Vehicle     67.0 ± 1.7     HA: 55.0     Mr auple     1/1       IA Vehicle     IA Vehicle     Cross, a new     2 ml of sterile     1/1       IA Vehicle     IA Vehicle     IA Avian (derived     NA	an of Treated perops/kge of perops/kge of 	Mof Treated         Origin of HA         Molecular         Dose         Number of participants         Follow-up creation         Conconitant OA           group/Age of metan ± SD or metan ± SD or ± SD or the S	motifiant P2-20         Constituent (mode)         Multice rol         Follow-up         Concontinuer (mode)         Multication           motifiant P2-20         weight         weight         cycles/muncher duration         cycles/muncher duration         consonitant (A)         Para provided type of AS/s           pristions         (weight         cycles/muncher duration         consonitant (A)         Para provided type of AS/s           pristions         (weither E2)         (or imperitons         (weither E2)         consonitant (A)         Para provided type of AS/s           pristions         (weither E2)         (or imperitons         (weither E2)         consonitant (A)         Para provided type of AS/s           pristions         (weither E2)         (or imperitons         (weither E2)         consonitant (A)         Para provided type of AS/s           pristions         (weither E2)         (or imperitons         (weither E2)         consonitant (A)         provided type of AS/s           pristions         (weither E2)         (or imperitons         (weither E2)         consonitant (A)         provided type of AS/s           pristions         (weither E2)         (or imperitons         (weither E2)         constroled thype of AS/s         consonitant (A)           pristions         (weither E2)         (or indiffigres)	Table 2 (continued)	Study Locatic OA	Dougados Knee 1993 [77]	Henderson Knee 1994 [78]	Henrotin Knee 2017 [79]	Heyworth Hand 2008 [80] (Basa joint thuml	Huskisson Knee 1999 [81]	<b>Jubb 2003</b> Knee [82]
Origin of HAMolecularDoseNumber of cycles/number of injections per cycleAvian500-730 kDaHyalectin 20 mg1/4extracted750,00020 mg Hyalagan in 1/5Avian500,0002 ml of sterile buffered saline1/1Avian500,0002 ml of sterile buffered saline1/1Avian500-730 kDa2 ml of sterile buffered saline1/1Avian500-730 kDa2 ml of sterile buffered1/5Avian500-730 kDa2 mg/2 ml1/5Avian500-730 kDa2 0mg/2 ml3/3Avian500-730 kDa2 0m	Origin of HAMolecularDoseNumber ofFollow-upweightweightDoseNumber ofFollow-upAvian500-730 kDaHyalectin 20 mg1/452 weeks(extracted from cock's500-730 kDaHyalectin 20 mg1/452 weeksAvian500-730 kDaHyalectin 20 mg1/452 weeks(extracted from cock's500-730 kDaHyalectin 20 mg1/450 weeksAvian500-700 Da2 mg (10 mg/ml)1/126 weeksAvian500-700 Da2 ml of sterile huffered saline1/126 weeksKartliage from roosterNANA1/11/126 weeksAvian (derived reticulated trom roosterNANA1/11/126 weeksAvian (derived from roosterNAS00-730 kDa20mg/Dml1/126 weeksAvian (derived from roosterNAS00-730 kDa20mg/Dml1/526 weeksAvian (derived from roosterS00-730 kDa20mg/Dml3/33/3S0 weeks	Origin of HA         Molecular         Dose         Number of cinjections         Follow-up cation)         Concontiant OA cation           Avian         500-730 kDa         Hyalectin 20 mg         1/4         52 weeks)         cation) allowed per cycle           Avian         500-730 kDa         Hyalectin 20 mg         1/4         52 weeks)         Any nonsteroidal matory drugs           Avian         500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal matory drugs           Avian         500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal anti-inflam- matory drugs           Avian         500-730 kDa         2 m of sterile         2 m of sterile         analgesia or anti- inflam- matory drugs           Avian         500-700 ba         2 m of sterile         2 m of sterile         analgesia or anti- inflam- matory           Avian         500-000         2 m of sterile         2 m of sterile         analgesia or anti- inflam- matory           NA         NA         2.2.ml, I6mg/ml         1/1         2 6 weeks         NSMDs. Topical drugs and viant derive           Ma         NA         1.4         2 6 weeks         NSMDs. Topical drugs are and viant derive         NSMDs. Topical drugs are and crossed           Aian         10	Origin of HA weight         Molecular before before the cycles/number duration per cycles         Number of Follow-up citypetions         Concontiant (AA         Data provided (red)         Data provided (red)           Avian         500-730 kDa         Hyalectin 20 mg Hya 20 mg/Hyalec		n of Treated groups/Age of participants (mean ± SD or median [P25- P75])	IA HA: 67 ± 9.7 IA Vehicle placebo: 69.0 ± 10.6	Data for sever- ity group 2. IA HA: 72.1 $\pm$ 1.7 IA Vehicle: 67.0 $\pm$ 1.7	IA HA: 66.9 ± 10.4 ± 8.5 ± 8.9	IAHA: 65.0 $1 \pm 2.0$ $\therefore$ IA SA: 64.0 $\Rightarrow 2.0$	IA HA: 65.8 ± 8.8 IA SA: 64.8 ± 9.3	IA HA: 63.5 ± 9.5 IA SA: 65.0 + 9.1
MolecularDoseNumber of cycles/number of injections per cycle500-730 kDaHyalectin 20 mg1/4500-730 kDaHyalectin 20 mg1/4500,000 Da20 mg Hyalgan in 1/51/5500,000 Da2 ml of sterile buffered saline1/1NA2 .2.ml, 16mg/ml1/1NA2 .2.ml, 16mg/ml1/1NA2 .2.ml, 16mg/ml1/1S00-730 kDa2 0mg/2ml1/5500-730 kDa2 0mg/2ml1/5500-730 kDa2 0mg/2ml3/3	MolecularDoseNumber ofFollow-upweightDoseNumber ofFollow-upweightcycles/numberduration500-730 kDaHyalectin 20 mg1/452 weeks)500-730 kDaHyalectin 20 mg1/452 weeks500,00020 mg Hyalgan in 1/522 weeks500,000 ba20 mg Hyalgan in 1/522 weeks500,000 ba20 mg Hyalgan in 1/526 weeks500,000 ba20 mg Hyalgan in 1/526 weeks500,000 ba20 mg Hyalgan in 1/526 weeks500,730 kDa20 mg/hyala1/126 weeks500,730 kDa20 mg/Dnl HA1/526 weeks500,730 kDa20 mg/Dnl HA3/352 weeks	Molecular         Dose         Number of cycles/number         Follow-up duration         Conconniant OA           weight         Dose         Number of cycles/number         Follow-up duration         Conconniant OA           500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal           500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal           500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal           500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Anti-inflam-           500-7500 ba         2 ml of sterile         22 weeks         Alternative           500000 ba         2 ml of sterile         anti-inflam-         anti-inflam-           NA         2.2ml, Iomg/ml         1/1         26 weeks         NSAIDs, Topical           NA         2.2ml, Iomg/ml         1/1         26 weeks         NSAIDs, Topical           NA         2.2ml         1/5         26 weeks         NSAIDs, Topical           S00-730 kDa         20mg/2ml         1/5         26 weeks         NSAIDs, Topical           S00-730 kDa         20mg/2ml         27         26 weeks         NSAIDs, Topical           S0	Molecular         Dose         Number of cilipications         Follow-up weight         Concomitant (A)         Data provided (ype of AE)% of patients           500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal         All AEs seem considered)           500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal         All AEs seem considered)           500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal         All AEs seem considered)           500-730 kDa         Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal         All AEs seem considered)           500.0000         20 mg Hyalgan in L/5         22 weeks         Allermative interoid during the considered         Summary/Not interoid during the considered           NA         2.2ml, I6mg/ml         1/1         26 weeks         NSAIDs, Topical         Summary of munbers of NSAIDs           NA         2.2ml, I6mg/ml         1/1         26 weeks         NSAIDs, Topical         Summary of munbers of NSAIDs         Summary of munbers of NSAIDs         All reported: interoid           NA         NA (1-mL injec- 1/2         26 weeks         NSAIDs, Topical         Summary Sub-730 kDa         Som230 kDa         Som230 kDa         Munbers of NSAIDs           500		Origin of HA	Avian (extracted from cock's combs)	Avian (extracted from rooster combs)	NA (Kartilage® Cross, a new reticulated HA supple- mented with mannitol)	Avian (derived from rooster combs)	Avian (extracted from rooster combs)	NA (Hyalgan <sup>®</sup> )
DoseNumber of cycles/number of injections per cycleHyalectin 20 mg1/4(10 mg/ml)1/420 mg Hyalgan in 1/522 ml of sterile buffered saline1/1HA1/1Somg/ml1/1101/120mg/ml1/120mg/zml1/520mg/zml1/520mg/zml3/3	DoseNumber of cycles/number duration of injections per cycleFollow-up duration duration (weeks) per cycleHyalectin 20 mg1/452 weeksL0 mg/ml)1/452 weeks20 mg Hyalgan in 1/522 weeks2 ml of sterile buffered saline22 weeks2 ml of sterile buffered saline1/126 weeks3 ml of sterile buffered saline1/126 weeks2 ml of sterile buffered saline1/33/33 ml of sterile buffered saline3/352 weeks	Dose     Number of cycles/number duration of injections     Follow-up treatment (medi- of injections     Concomitant OA treatment (medi- nation) allowed       Hyalectin 20 mg     14     52 weeks     Any nonsteroidal nummer duration       Hyalectin 20 mg     14     52 weeks     Any nonsteroidal nummer duration       20 mg Hyalgan in 1/5     22 weeks     Any nonsteroidal or analgesics       20 mg Hyalgan in 1/5     22 weeks     Alternative analgesics or anti- inflammatory treatment (per- mited during the follow up but of during the treat- ment period)       2.2ml, 16mg/ml     1/1     26 weeks     NSAIDs, Topical during the treat- ment period)       NA (1-mL injec-     1/2     26 weeks     NSAIDs, Groung during the treat- ment period)       20mg/2ml     1/5     26 weeks     NSAIDs, Topical during the treat- ment period)       20mg/2ml     1/5     26 weeks     NSAIDs, Groung during the treat- ment period)       20mg/2ml     3/3     52 weeks     NSAIDS, Sibu- secetion       20mg/2ml     3/3     52 weeks     Rice use of indicates and indicates	Dose         Number of cycles/number of injections         Follow-up (veeks)         Conconnitant OA         Data provided in the article of patients           Hyalectin 20 mg         1/4         52 weeks         atti-inflam- (NSAIDs) and/ of sterile         Any nonsteroidal         All AEs seen individues           (10 mg/ml)         1/4         52 weeks         Any nonsteroidal         All AEs seen individues           20 mg Hyalectin 20 mg         1/4         52 weeks         Any nonsteroidal         All AEs seen individues           20 mg Hyalegan in 1/5         22 weeks         Any nonsteroidal         All AEs seen inflam- inflammatory         the article           2 ml of sterile         1/5         22 weeks         Alternative         SummatyNot           2 ml of sterile         NA (1-mL injec-         1/1         26 weeks         NSAIDs, Topical           NA (1-mL injec-         1/5         26 weeks         NSAIDs, Topical         All reported: inflammatory           20mg/2ml         1/5         26 weeks         NSAIDs, Topical         Mireported: indiamatory           20mg/2ml         3/3         52 weeks         NSAIDs (ibu- initia mative         All reported: indiamatory           20mg/2ml         1/5         26 weeks         NSAIDs (ibu- initia mative         All reported: indiamatory           Song		Molecular weight	500-730 kDa	500,000 -750,000 Da	Ч Ч	NA	500–730 kDa	500–730 kDa
Number of cycles/number of injections per cycle 1/4 1/2 1/2 1/5 3/3	Number of cycles/number duration of injections lldFollow-up duration duration1/452 weeks1/422 weeks1/126 weeks1/226 weeks1/526 weeks3/352 weeks	Number of evcles/number of injectionsFollow-up treatment (medi- cation) allowed per cycle1/452 weeksAny nonsteroidal anti-inflam- matory drugs (NSAIDs) and/ or analgesics1/452 weeksAny nonsteroidal anti-inflam- matory drugs inflammatory treatment (per- mitted during the follow up but not during the reat- ment period)1/226 weeksNSAIDs, Topical NSAIDs1/526 weeksNSAIDs, Topical not every 4-6 h, as needed)3/352 weeksNSAIDs (jbu- profen 400 mg every 4-6 h, as nof noreded)3/352 weeksNSAIDs (stron- noreded)3/352 weeksNSAIDs except indometorion	Number of cycles/number duration tereatment (medi- of injections (weeks)     Concomitant OA cation) allowed (type of AE% of Patients considered)       1/4     52 weeks     Any nonsteroidal     All AEs seem anti-inflam- natory drugs       1/4     52 weeks     Any nonsteroidal     All AEs seem anti-inflam- or analgesics       1/15     22 weeks     Any nonsteroidal     All AEs seem anti-inflam- or analgesics       1/1     26 weeks     NSAIDS) and/ or analgesics     Summary/Not analgesics       1/1     26 weeks     NSAIDS, Topical     Summary of AEs       1/2     26 weeks     NSAIDS, Topical     Summary of AEs       1/3     26 weeks     NSAIDS, Topical allowed observed dur- ing the study"       1/3     50 weeks     NSAIDS     Summary of AEs       1/3     52 weeks     NSAIDS     Summary of pocal reac- tions (in the majority of pocal reac- tions (in the post common pocal reac- tions (post common post commo		Dose	Hyalectin 20 mg (10 mg/ml)	20 mg Hyalgan i 2 ml of sterile buffered saline	2.2ml, 16mg/ml HA	NA (1-mL injec- tion of hylan G-F 20)	20mg/2ml	20mg/2ml HA
	Follow-up         • duration         (weeks)         52 weeks         22 weeks         26 weeks	Follow-upConcomitant OAdurationtreatment (medi-(weeks)cation) allowed52 weeksAny nonsteroidalanti-inflam-matory drugsNSAIDsNSAIDs and/or analgesicsanti-inflam-and or analgesicsanti-inflam-22 weeksAlternativeanalgesicsanti-inflam-profenduring the treat-inflammatory26 weeksNSAIDs, Topical26 weeksNSAIDs (ibu-profen 400 mgevery 4-6 h, as26 weeksNSAIDs (ibu-profen 400 mgevery 4-6 h, as25 weeksNSAIDs (ibu-52 weeksNSAIDs (ibu-free use ofanalgesics andanalgesicsanalgesics andMonethorinindomethorion	Follow-up duration       Concomitant OA       Data provided in the article (type of AE/% of patients         52 weeks       Any nonsteroidal       All AEs seem anti-inflam- to have been matory drugs         52 weeks       Any nonsteroidal       All AEs seem anti-inflam- to have been matory drugs         22 weeks       Any nonsteroidal       All AEs seem anti-inflam- natory drugs         23 weeks       Any nonsteroidal       All AEs seem anti-inflam- netory drugs         26 weeks       NSAIDs, Topical       Summary of numbers of AEs         26 weeks       NSAIDs, Topical       Summary of numbers of AEs         26 weeks       NSAIDs       All reported: "No AEs         26 weeks       NSAIDs       AIL reported: "No AE         26 weeks       NSAIDs       AIL reported: "No AE         26 weeks       NSAIDs       Summary of numbers of AEs         26 weeks       NSAIDs       Summary of numbers of AEs         27 weeks       NSAIDs       Summary focusing in local reac- tifor reactions         27 weeks       Stee veroid AE       AE         28 weeks       Stee veroid AE       AE         28 weeks       NSAIDs except       AE         AE       AE       AE         AE       AE       AE         AE       AE		Number of cycles/number of injections per cycle	1/4	n 1/5	1/1	- 1/2	1/5	3/3
Concomitant OA       Data provided in the article       Published data usable (type of AE/% for M-A?)         cation) allowed       (type of AE/% for M-A?)       (type of AE/% for M-A?)         Any nonsteroidal       All AEs seem       Yes/no)         Any nonsteroidal       All AEs seem       Yes         anti-inflam-       to have been       Yes/no)         matory drugs       reported       (yes/no)         or analgesics       Summary/Not       No         Alternative       Summary/Not       No         analgesics or anti- follow up but not during the follow up but not during the treat- ment period)       No         NSAIDs, Topical       Summary of numbers of AEs       No         NSAIDs (ibu- meteded)       All reported:       Yes         NSAIDs (ibu- meteded)       No focusing in local reac- tions (in the       No         Sammary       No       No       No         Summary       No       No <td>Data provided in the article (type of AE/% for M-A? of patients (ves/no)Published data usable (ves/no)(type of AE/% for M-A? or M-A? considered)Yes (ves/no)All AEs seem reportedYesAll AEs seem detailedYesSummary of detailedNo detailedAll reported: mumbers of hes observed dur- ing the study"YesSummary observed dur- ing the study"No focusing in focusing in<b< td=""><td>Published data usable for M-A? (yes/no) No Yes Yes Yes</td><td></td><td></td><td>Full data provided by the author/sponsor? (source of informa- tion)</td><td>Author contacted with no response</td><td>No contact informa- tion found</td><td>No (Laboratories VIVACY, France. Data promised but not provided.)</td><td>Author contacted with no response</td><td>No contact informa- tion found</td><td>No (Fidia Pharma)</td></b<></td>	Data provided in the article (type of AE/% for M-A? of patients (ves/no)Published data usable (ves/no)(type of AE/% for M-A? or M-A? considered)Yes (ves/no)All AEs seem reportedYesAll AEs seem detailedYesSummary of detailedNo detailedAll reported: mumbers of hes observed dur- ing the study"YesSummary observed dur- ing the study"No focusing in focusing in <b< td=""><td>Published data usable for M-A? (yes/no) No Yes Yes Yes</td><td></td><td></td><td>Full data provided by the author/sponsor? (source of informa- tion)</td><td>Author contacted with no response</td><td>No contact informa- tion found</td><td>No (Laboratories VIVACY, France. Data promised but not provided.)</td><td>Author contacted with no response</td><td>No contact informa- tion found</td><td>No (Fidia Pharma)</td></b<>	Published data usable for M-A? (yes/no) No Yes Yes Yes			Full data provided by the author/sponsor? (source of informa- tion)	Author contacted with no response	No contact informa- tion found	No (Laboratories VIVACY, France. Data promised but not provided.)	Author contacted with no response	No contact informa- tion found	No (Fidia Pharma)

스 Adis

Table 2 (cont	tinued)										
Study	Location of OA	Treated groups/Age of participants (mean ± SD or median [P25- P75])	Origin of HA	Molecular weight	Dose	Number of cycles/number of injections per cycle	Follow-up duration (weeks)	Concomitant OA treatment (medi- cation) allowed	Data provided in the article (type of AE/% of patients considered)	Published data usable for M-A? (yes/no)	Full data provided by the author/sponsor? (source of informa- tion)
Kwon 2013 [83]	Gleno- humeral joint (shoulder)	IA HA: 66.1 ± 10.7 IA SA: 66.1 ± ) 11.7	Avian (extracted from rooster combs)	620,000- 1,170,000 E	NA Da	1/3	26 weeks	Current regimen of pain medica- tions could be maintained but no additional treatment to the shoulder	Report mainly focused on device related AEs. AEs by SOC not reported, but "Serious" AEs reported	Yes	Author contacted with no response
Lohmander 1996 [84]	Knee	IA HA: 58.53 ±8.34 IA SA: 58.03 ±8.44	NA (Hyaluro- nan, Artzal®)	1000 kDa (Average)	25mg/2.5ml	1/5	20 weeks	Simple analgesics, in addition to NSAIDs	Summary/Not detailed	No	No (Author)
Lundsgaard 2008 [85]	Knee	IA HA: 68.8 ± 6.27 6.27 IA SA: 69.6 ± 7.27	NA (Hya- luronate, Hyalgan <sup>®</sup> )	A	2ml of Hyalgan, 10.3 mg/ml	1/4	26 weeks	NSAID (inclusive COX-2 selec- tive inhibitors), codeine, and tramadol	No AE (serious non-serious, local reaction, post injec- tion 'flares') reported during the trial. But 1 withdrawal after 2 weeks due to cerebra haemorrhage	Yes	Author contacted with no response
Munteanu 2011 [86]	First MTPJ (foot)	IA HA: 53.7 ± 11.3 IA SA: 55.3 ± 11.2	NA (Hylan G-F 20, Synvisc)	NA	NA (up to 1 ml of hylan G-F 20)	<ul> <li>1/1 (with an option</li> <li>of a 2nd and final injection</li> <li>at month 1 or</li> <li>3 if there was</li> <li>no improvement)</li> </ul>	26 weeks	Not clear: "Use of (paraceta- mol rescue medication) and co-interventions to relieve first MTPJ pain, as secondary outcome"	Summary/Not detailed	ç	Author contacted with no response
Navarro- Sarabia 2011 [87]	Knee	IA HA: 63.0 ± 8.2 IA SA: 63.9 ± 8.9	Obtained from strains of Streptococ- cus zoopi- demicus	900,000 Da (Average)	2.5 ml 1% sodium hyalu- ronate	4/5	174 weeks	Short cycles of NSAID	Numbers of AEs reported, not per SOC frequencies	No	Author contacted with no response

If TreatedOrigin of HAMolecularDoseNumber of cycles/numbergroups/Age of groups/Age of metaia [P25-Origin of HAMolecularDoseNumber of cycles/number $P51$ ) $Rean \pm SD$ or metaia [P25- $Rean \pm SD$ or metaia [P25- $Rean \pm SD$ or metaia [P25- $Rean \pm SD$ or sold the hA, LMW: $NA$ $1/3$ $IA$ HA, LMW: $AHA$ , LMW: $AHA$ , LMW: $AHA$ , LMW: $AHA$ , LMW: $I/3$ $IA$ HA, LMW: $B00 \pm 5.0$ $B00 \pm 5.0$ $HMW$ : $HMW$ : $I/3$ $IA$ HA, LMW: $B00 \pm 5.0$ $HMW$ : $I/12$ to $2.0$ $I/3$ $IA$ HA, LMW: $IAA$ , LMW: $IAA$ , LMW: $I/3$ $I/3$ $IA$ HA, LMW: $IAA$ , LMW: $IAA$ , LMW: $I/3$ $I/3$ $IA$ HA, LMW: $IAA$ , LMW: $IAA$ , LMW: $I/3$ $I/3$ $IAA$ , LMW: $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , EGO $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $IAA$ , ESO $I/3$ $IAA$ , ESO	$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	of Treated participantsOrigin of HA supportMolecular weightDose cycles/mumber duration origisationsConconitant OA caration attaininggrup/sgond mean $\pm$ SD or rotein P75])MAL DMW: NADowe origisationsConconitant OA caration origisationsConconitant OA caration process)Conconitant OA caration processConconitant OA processConconitant OA processConconitant OA caration processConconitant OA processConconitant OA processConconitant OA processConconitant OA proce		Table 2 (continued)	Study Location o OA	Petrella 2008 Knee [88]	Pham 2004 Knee [89]	Qvistgaard Hip 2006 [90]	Richette Hip 2009 [91]	Strand 2012 Knee [92]
Origin of HAMolecularDoseNumber of cycles/numberNAweightboseNumber of cycles/numberNAS80-780 kDa+1.2 to 2.0nillion Da+1.2 to 2.0HMW:580-730 kDaHMW:6 mil- lion DaStreptococ- cus equus1.900 kDaStreptococ- cus equus1.900 kDaStreptococ- cus equus2.5 mj of sodiumNANANANAStreptococ- cus equus00,000 DaMA900,000 DaStreptococ- cus fermen- tationNA (2.5 ml of HA)AvianNA<	Origin of HAMolecularDoseNumber ofFollow-upweightboseNumber ofFollow-upweightboseNumber ofFollow-upNADMW:NA1/316 weeksNaDMW:NA1/316 weeksS00-730 kDa1/12 to 2.0million Da1/316 weeksFuduced from1.900 kDa25 mg of sodium3/352 weeksSreptocor-1.900 kDa25 mg of sodium3/352 weeksNANANANA1/313 weeksProduced from1.900 kDa25 mg of sodium3/352 weeksStreptocor-2.5 ml1/31/313 weeksUhalgan®NANANA1/313 weeksStreptocor-cus fermen-2.5 ml1/313 weeksVialion900,000 DaNA (2.5 ml of1/113 weeksStreptocor-sudNANA1/313 weeksStreptocor-sudNA (2.5 ml of1/113 weeksStreptocor-sudNA3.0 mg HA in1/113 weeksStreptocor-sudNA3.0 mg HA in1/113 weeks	Origin of HA         Molecular         Dose         Number of cycles/number duration per cycle         Follow-up treament (medi- of injections         Concomitant OA           NA         DMW:         0f injections         (weeks)         cation) allowed of injections         cation) allowed           NA         DMW:         MA         I/3         16 weeks         Analgesics           NA         DMW:         6 million Da allion Da 1000 Da         1/3         16 weeks         Analgesics           NA         DMW:         6 million Da allion Da         1/3         16 weeks         Analgesics           Streptoctor         1000 kba         25 mg of sodium         3/3         52 weeks         NSAIDs           Streptoctor         1.900 kba         1/3         1/3         13 weeks         Usual analgesic consumption (flyalgan <sup>6</sup> )           Obtained from cus equus         00.000 Da         NA         1/3         1/3         13 weeks         NSAIDs           Streptococ- cus fermen- tution         90.000 Da         NA         1/3         1/3         13 weeks         NSAIDs, herhal (with no other information)           Main         Ma         1/1         13 weeks         NSAIDs, herhal (with no other information)         Analgesics           Main         Ma         3.01	Origin of HA         Molecular         Dose         Number of cycles/minter         Follow-up conjunction         Conconitant OA         Data provided cycles/minter           NA         Molecular         Dose         Number of cycles         Follow-up cycles         Conconitant OA         Data provided cycles/minter           NA         DMW:         MA         1/3         16 weeks)         cation) allowed         in the article considered)           NA         DMW:         MA         1/3         16 weeks         Anallesics         catalid cycles/minter           NM         S80-780 kDa million Da S80-730 kDa 1,000 La         NA         1/3         16 weeks         Anallesics         Soumary/Not           Produced from infinen Da Sweptoco- cus equats         1.000 kDa         2.5 ml         Soumary         Soumary/Not           Produced from information         1.900 kDa         2.5 ml         13 weeks         NSAIDs)         Source some source dust vorter out or other         Source some source dust vorter out other         Source some source dust vorter out other         Source some source dust vorter out other         Source some vorter out other         Anallesis or ofp out other         Source some vorter out other           NA         NA         NA         NA         NA         NA         NA         Source sop vorter out other         Ana		of Treated groups/Age of participants (mean ± SD or median [P25- P75])	IA HA, DMW: 68.0 $\pm$ 6.0 1A HA, LMW: 69.0 $\pm$ 5.0 1A HA, HMW: 71.0 $\pm$ 9.0 IA SA: 71 $\pm$ 8	IA HA: 64.9 ± 8.4 IA SA: 64.9 ± 7.7	IA HA: 65.0 ± 14.0 IA SA: 64.0 ± 11.0	IA HA: 60.8 ± 10.2 IA SA: 59.5 ± 12.6	IA HA: 60.9 ± 10.24 IA SA: 60.3 ± 9.97
MolecularDoseNumber of cycles/numberweightDoseNumber of cycles/numbermeightof injections of injectionsDMW:NA1/3580-780 kDa1/3+ 1.2 to 2.0million DaLMW:55 mg of sodium500-730 kDa3/3hyaluronate in 2.5 ml3/3900,000 DaNA (2.1 HA)900,000 DaNA (2.5 ml of HA)NA30 mg HA inNA30 mg HA in	MolecularDoseNumber ofFollow-upweightDoseNumber durationweightcycles/number durationper cycle(weeks)S80-780 kDa1/316 weeks580-730 kDa1/316 weeksh112 to 2.0million Da3/352 weeksh12 to 2.0hyaluronate in3/352 weeksh12 to 2.01/31/313 weekss80-730 kDa25 mg of sodium3/352 weekshMW: 6 mil-2.5 ml1/313 weekshMW: 6 mil-1/31/313 weekshMW: 6 mil-1/31/313 weeksbion DaNA1/313 weekshMW: 6 mil-1/31/313 weekshMW: 6 mil-1/31/313 weekshMW: 6 mil-1/31/313 weekshMW: 6 mil-1/113 weeks1/3hMW: 6 mil-1/113 weekshMW: 6 mil-1/113 weekshMM: 73 mg HA in1/1hM3 mg HA in1/1hM3 mg HA in1/1hM1/113 weeks	Molecular         Dose         Number of cycles/humber of injections         Follow-up (weeks)         Conconitant OA retation           MW:         NA         1/3         16 weeks         ation) allowed per cycle           DMW:         NA         1/3         16 weeks         Analgesics           \$80-780 kba         NA         1/3         16 weeks         Analgesics           \$80-790 kba         NA         1/3         16 weeks         Naalgesics           \$80-790 kba         1/3         16 weeks         Naalgesics           \$80-790 kba         1/3         16 weeks         NSAIDs)           million Da         S00-730 kba         NSAIDs)         NSAIDs)           MW:         6 mil- lion Da         NSAIDs         NSAIDs)           MOKDa         25 mg of sodium         3/3         52 weeks         NSAIDs)           900,000 ba         NA         1/3         13 weeks         Usual analgesic (with no other information)           900,000 ba         NA (2.5 ml of HA)         1/1         13 weeks         NSAIDs, nered information)           900,000 ba         NA (2.5 ml of HA)         1/1         13 weeks         NSAIDs, nered information)	Molecular         Dose         Number of cycles/number duration         Follow-up terment (medi- per cycles         Concomitant OA         Data provided (type of AE)%           Molecular         Dose         Number of cycles/number         Follow-up cycles/number         Concomitant OA         Data provided (type of AE)%           DMW:         NA         1/3         16 weeks)         cation) allowed         (type of AE)%           St0-780 kDa         1/3         16 weeks         Analgesics         considered)           DMW:         NA         1/3         16 weeks         Analgesics         considered)           DMW:         Analgesics         Summary/Not         considered)         considered)         considered)           DMW:         Analgesics         Summary/Not         considered)         considered)           DMW:         Analgesics         Summary/Not         considered)         considered)           DMW:         Analgesics         Son stoped         considered)         considered)           DMW:         Analgesics         Son stoped         considered)         considered)           DMW:         Analgesics         Son stoped         considered         considered           DMW:         Analgesics         Son stoped         considered		Origin of HA	YN	Produced from Streptococ- cus equus	: NA (Hyalgan <sup>®</sup> )	Dobtained from Streptococ- cus fermen- tation	: Avian (Chicken combs)
DoseNumber of cycles/numberDoseNumber of cycles/numberNA1/3NA1/3Da3/3hyaluronate in 2.5 ml3/3NA1/3NA1/3NA1/33 ml1/1	Dose     Number of cycles/number duration of injections     Follow-up duration       NA     1/3     16 weeks       NA     1/3     16 weeks       Da     1/3     13 weeks       NA     1/3     13 weeks       NA     1/1     13 weeks       NA     1/1     13 weeks       MA     1/1     13 weeks       MA     1/1     13 weeks	DoseNumber of cycles/number durationFollow-up treatment (medi- of injections weeks)Concomitant OA reatment (medi- of injections of injections analgesics (including NSAIDs)NA1/316 weeksAnalgesics (including NSAIDs)NA1/316 weeksAnalgesics (including NSAIDs)Da1/316 weeksNalgesics (including NSAIDs)Da1/316 weeksNSAIDs)Da25 mg of sodium3/352 weeksNSAIDs)Da2.5 ml1/313 weeksNSAIDs)Da1/313 weeksUsual analgesic consumption (with no other information)NA1/113 weeksNSAIDs or step 2 analgesics oral therapies, oral HA, glucosa- mine, chon-	Dose     Number of cycles/number of injections     Follow-up (weeks)     Conconitant OA     Data provided in the article of patients       NA     1/3     16 weeks     Analgesics     SummaryNot       NA     1/3     16 weeks     Analgesics     SummaryNot       NA     1/3     16 weeks     NSAIDs)     SommaryNot       NA     1/3     16 weeks     NSAIDs)     SommaryNot       Sa     1/3     16 weeks     NSAIDs)     SommaryNot       Data     NA     1/3     13 weeks     NSAIDs)     Sof patients       Data     NA     1/3     13 weeks     NSAIDs)     Sof crepating       Data     NA     1/3     13 weeks     NSAIDs)     Sof crepating       NA     1/3     13 weeks     Usual analgesic     SummaryNot       NA     1/1     13 weeks     Usual analgesic     SummaryNot       MA     1/1     13 weeks     NSAIDs or step 2     SummaryNot       MA     1/1     13 weeks     NSAIDs or step 2     SummaryNot       MA     1/1     13 weeks     NSAIDs or step 2     SummaryNot       MA     1/1     13 weeks     NSAIDs or step 2     SummaryNot       MA     1/1     13 weeks     NSAIDs or step 2     SummaryNot		Molecular weight	DMW: 580-780 kT 580-780 kT + 1.2 to 2.0 million Da 1LMW: 500-730 kT HMW: 6 mil- lion Da	1.900 kDa	AN	900,000 Da	ΥN
Number of cycles/number of injections per cycle 1/3 1/3 1/1 1/1	Number of cycles/number duration of injections l/3Follow-up duration duration get cycle1/316 weeks1/316 weeks1/313 weeks1/113 weeks1/113 weeks	Number of cycles/number duration of injections per cycle     Follow-up duration treatment (medi- cation) allowed       1/3     16 weeks     Analgesics (including NSAIDs)       1/3     16 weeks     Analgesics (including NSAIDs)       1/3     16 weeks     NSAIDs)       1/3     13 weeks     NSAIDs)       1/3     13 weeks     NSAIDs)       1/1     13 weeks     NSAIDs or step 2 analgesics       1/1     13 weeks     NSAIDs, herbal therapies, oral HA, glucosa- min, chon-	Number of cycles/number of injections per cycle     Follow-up treatment (medi- for her article of patients considered)       1/3     16 weeks     Analgesics considered)       1/3     16 weeks     Analgesics considered)       1/3     16 weeks     Analgesics considered)       1/3     16 weeks     NSAIDs)       1/3     52 weeks     NSAIDs)       1/3     52 weeks     NSAIDs)       1/3     13/3     52 weeks       1/3     13 weeks     NSAIDs)       1/1     13 weeks     NSAIDs)       1/1     13 weeks     Usual analgesic consumption       1/1     13 weeks     Usual analgesic consumption       1/1     13 weeks     NSAIDs or step 2       1/1     13 weeks     NSAIDs, herbal       1/1     13 weeks     NSAIDs, herbal       1/1     13 weeks     NSAIDs, or step 2       1/1     13 weeks     NSAIDs, herbal       1/1     13 weeks     NSAIDs, or step 2       1/1     13 weeks     NSAIDs, or step 2       1/1     13 weeks     NSAIDs		Dose	Da Da	25 mg of sodiun hyaluronate in 2.5 ml	NA (2 ml HA)	NA (2.5 ml of HA)	30 mg HA in 3 ml
	Follow-up t duration (weeks) 16 weeks 52 weeks 13 weeks 13 weeks 13 weeks	Follow-up       Concomitant OA         r duration       treatment (medi- cation) allowed         16 weeks       Analgesics (including NSAIDs)         52 weeks       NSAIDs)         13 weeks       Usual analgesics (with no other information)         13 weeks       NSAIDs or step 2 analgesics         13 weeks       NSAIDs, herbal         13 weeks       NSAIDs, or step 2 analgesics         13 weeks       NSAIDs, or step 2 analgesics         13 weeks       NSAIDs, herbal         13 weeks       NSAIDs, or step 2 analgesics         13 weeks       NSAIDs, herbal         HA, glucosa- mine, chon-       Interapies, oral	Follow-up (weeks)       Concomitant OA cation) allowed       Data provided (type of AE/% of patients         16 weeks       Analgesics       Summary/Not         16 weeks       NSAIDs)       Most common!         52 weeks       NSAIDs)       Most common!         13 weeks       Usual analgesics       Summary/Not detailed         13 weeks       NSAIDs or step       Sonc specific AEs for GI         13 weeks       NSAIDs or step       Sommary/Not detailed         13 weeks       NSAIDs or step       Summary/Not detailed         13 weeks       NSAIDs, herbal       Focus on theraties, oral         13 weeks       NSAIDs, herbal       Focus on theraties, o		Number of cycles/number of injections per cycle	1/3	л 3/3	1/3	T/I	1/1
Concomitant OAData providedPublishedtreatment (medi-in the articledata usablecation) allowed(type of AE/%for M-A?(type of AE/%for M-A?of patients(yes/no)of patientsof patients(yes/no)considered)AnalgesicsSummary/NotNo(includingdetailedNSAIDsMost commonly Yesseen AEsreported. BySOC reportfor someSOCs, andonly specificUsual analgesicsSummary/NotNoconsumption(with no otherinformation)AEs for GINoNSAIDsNSAIDs or step 2Summary/NotNoNSAIDs or step 2Summary/NotNoanalgesicsAll AEs notfor bablyhth, glucosa-SOC resultsAll AEs notHA, glucosa-SOC resultsfor someinformation)AEs reported.Soc resultsinformation)AEs reported.for someSome specific(probablyfreatment)AEs reported.SoC resultsinformationAll AEs notinformationfor someinformationSoC resultsinformationfor someinformationfor someinformationfor someinformationfor someinformationfor someinformationfor someinformationfor someinformationfor someinformationfor someinformationfor some </td <td>Data provided       Published         In the article       data usable         (type of AE/%       for M-A?         (type of AE/%       for M-A?         of patients       (yes/no)         considered)       yes/no)         Summary/Not       No         detailed       No         Most commonly Yes       seen AEs         reported. By       SOCs, and         SOCS, and       No         adtailed       No         AEs for GI       Summary/Not         All AEs not       AEs for GI         Some specific       (probably         for bably       AEs reported.         Some specific       (probably         AEs reported.       Some specific         Some specific       (probably         for results       No         AEs reported.       Yes         Some specific       Sources unted;         AEs reported.       Yes         Focus on       Yes</td> <td>Published data usable for M-A? (yes/no) No No Yes</td> <td></td> <td></td> <td>Full data provided by the author/sponsor? (source of informa- tion)</td> <td>Author contacted with no response</td> <td>Author contacted with no response</td> <td>No (Fidia Pharma)</td> <td>No (Author)</td> <td>Author contacted with no response</td>	Data provided       Published         In the article       data usable         (type of AE/%       for M-A?         (type of AE/%       for M-A?         of patients       (yes/no)         considered)       yes/no)         Summary/Not       No         detailed       No         Most commonly Yes       seen AEs         reported. By       SOCs, and         SOCS, and       No         adtailed       No         AEs for GI       Summary/Not         All AEs not       AEs for GI         Some specific       (probably         for bably       AEs reported.         Some specific       (probably         AEs reported.       Some specific         Some specific       (probably         for results       No         AEs reported.       Yes         Some specific       Sources unted;         AEs reported.       Yes         Focus on       Yes	Published data usable for M-A? (yes/no) No No Yes			Full data provided by the author/sponsor? (source of informa- tion)	Author contacted with no response	Author contacted with no response	No (Fidia Pharma)	No (Author)	Author contacted with no response

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Table 2 (con	tinued)										
Study	Location o OA	f Treated groups/Age of participants (mean ± SD o median [P25- P75])	Origin of HA	Molecular weight	Dose	Number of cycles/number of injections per cycle	Follow-up duration (weeks)	Concomitant OA treatment (medi- cation) allowed	Data provided in the article (type of AE/% of patients considered)	Published data usable for M-A? (yes/no)	Full data provided by the author/sponsor? (source of informa- tion)
Wobig 1998 [93]	Knee	IA HA: 60.0 ± 2.0 IA SA: 64.0 ± 2.0	NA (Hylan G-F 20)	NA	NA (2 ml of Hylan G-F 20)	1/3	26 weeks	Steroids, NSAIDs, analgesics, or any other therapy for the treatment of OA	Summary/Not detailed	No	No contact informa- tion found
Where publis sor	hed data wer	e adequate for in	clusion in the M-	-A and a full s	ufety report was also	provided by the	e author/spo	nsor, we preferentia	lly used the full	data obtained	from the author/spon-
AE adverse e	vent, COX-2	cyclooxygenase	2, DMW dual m	nolecular weigł	it, GI gastrointestir	nal, HA hyaluroi	nic acid, HM	W high molecular	weight, IA intra	-articular, IAI	44 intra-articular hya-

uronic acid. IASA intra-articular saline, LMW low molecular weight, M-A meta-analysis, MTPJ metatarsophalangeal joint, NA not available (i.e. information not provided in the manuscript)

VSAID non-steroidal anti-inflammatory drug, OA osteoarthritis, P25–P75 25th percentile–75th percentile, SD standard deviation, SOC System

the findings on quality assessment for all outcomes assessed in this meta-analysis, showing the overall analysis data. The certainty of the evidence was the same overall and with "studies without any concomitant anti-OA medication" for almost all the outcomes, apart from "severe adverse events" for which quality was graded as "low" (in contrast to "moderate", overall) (data not shown). When considering the studies with concomitant anti-OA medication, the certainty of the evidence for "nervous system disorders" was rather "low" (in contrast to "moderate" for the other groups), and was "moderate" for "vascular disorders" and "hypersensitivity reaction" (in contrast to "low" for the other groups) (data not shown). These differences in the quality of evidence were due to differences in imprecision around the estimates.

## 4 Discussion

Overall this meta-analysis found no increased odds of total AEs with IAHA compared with placebo; this is particularly true for studies without concomitant anti-OA medication (OR = 1.19, 95% CI 0.92–1.54), but also for studies that allowed concomitant anti-OA medication. When considering only the studies which responded to our selection criteria, particularly the criteria related to the non-use of concomitant anti-OA medications during the trials, we found no statistically significant increased odds of SAEs with IAHA compared to placebo, even though there are more SAEs with IAHA (OR = 1.78, 95% CI 0.92–3.47).

However, we found significant increased odds of SAEs in the IAHA group versus placebo, overall and particularly in studies with concomitant OA treatment allowed (OR = 1.78, 95% CI 1.10-2.89). This compares with the findings of a meta-analysis from Rutjes et al., which found a 41% increased relative risk (RR) of SAEs (RR = 1.41, 95%CI 1.01–1.74) [33]. In our analysis, the main studies leading to these results are the Jubb et al. 2003 study [82], the Kwon et al. 2013 study [83], and the Strand et al. 2012 study [92], respectively counting for 33.29, 13.03, and 1.87% of the total weight of all studies. In the Jubb et al. 2003 study, the free use of analgesics and NSAIDs except indomethacin was permitted during the trial [82]. In the Kwon et al. 2013 study, the usual regimen of pain medications could be maintained, but no additional treatment to the shoulder was allowed during the trial [83]. In Strand et al. [92] study, NSAIDs, herbal therapies, oral HA, glucosamine, chondroitin sulfate, minocycline and short-acting oral opiates were allowed during the trial. Oral NSAIDs are associated with an increased risk of SAEs [94], and the increased odds of SAEs reported in these studies might be due to the concomitant use of NSAIDs or other medications. These results are, however, difficult to interpret due to the paucity in the reporting of safety data for IAHA.



# IAHA: Infections and infestations

Fig. 4 Forest plot displaying the results of the meta-analyses comparing infections and infestations with IAHA versus placebo in patients with OA: analysis on studies without concomitant anti-OA medica-

tion allowed, analysis on studies with concomitant anti-OA medication allowed, and overall analysis. *CI* confidence interval, *IAHA* intraarticular hyaluronic acid, *OA* osteoarthritis

Like the Rutjes meta-analysis [33], our analysis includes data from the trial of Strand et al. of a cross-linked HA product (Gel-200) [92]. While the trial found a similar rate of AEs between IAHA and saline placebo, eight cases of SAEs were reported in the Gel-200 group, all judged unrelated to study treatment, including five cancers diagnosed soon after treatment administration. The biological plausibility of a link between IAHA and the SAEs reported has been questioned in the literature [95, 96], and there is no pre-clinical data to suggest any carcinogenicity with Gel-200 [92]. In contrast with our meta-analysis, which includes all types of HA, two meta-analyses of US-approved HA products have found no statistically significant differences between IAHA and IA placebo for any safety outcomes [34, 35].

We found a significant increase in odds of withdrawals due to AEs associated with IAHA overall (+ 62%; 95% CI

1.04–2.51), which was particularly high in studies of IAHA without concomitant anti-OA treatment (+ 80%) although this result did not reach significance (95% CI 0.99–3.26). This is in agreement with the Rutjes meta-analysis, which did not make any difference in the studies regarding to the use of concomitant anti-OA medications and also found an overall increased risk of dropouts due to AEs (RR = 1.33; 95% CI 1.01–1.74) [33].

We observed a high rate of selective outcome reporting in the studies included in this meta-analysis; over 50% of studies did not adequately report safety data (Fig. 3a, b). Only two authors (Jorgensen et al. [65]; van der Weegen et al. [72]) shared full safety reports with us for the purpose of this meta-analysis, and no full safety report was obtained for any of the studies with concomitant anti-OA medications. This may have led to a very large underestimation of

# Intra-articular hyaluronic acid: Serious adverse events

Study	n Active	N Active	n Placebo	N Placebo				Odds Ratio (95% Cl	% ) Weight
Without concomitan	t OA treat	ment							
Altman 2004	7	173	3	174				2.40 (0.61, 9.45)	8.14
Brandt 2001	6	114	4	112	_			1.50 (0.41, 5.47)	9.13
Jorgensen 2010	10	165	7	170		<b>-</b>		1.50 (0.56, 4.05)	15.56
Puhl 1993	2	102	0	107				5.35 (0.25, 112.76)	1.64
Altman 1998	0	164	0	168				(Excluded)	0.00
Baltzer 2009	0	135	0	107				(Excluded)	0.00
Dixon 1988	0	30	0	33				(Excluded)	0.00
Salk 2006	0	9	0	8				(Excluded)	0.00
Van der Weegen 20	150	99	0	97				(Excluded)	0.00
Subtotal (I-squared	= 0.0%, p	o = 0.83	4)			$\triangleleft$		1.78 (0.92, 3.47)	34.47
						Ť			
With concomitant O	A treatme	nt							
Altman 2009	9	293	9	295	_			1.01 (0.39, 2.57)	17.34
Jubb 2003	27	208	14	200				1.98 (1.01, 3.90)	33.29
Kwon 2013	11	150	5	150			-	2.29 (0.78, 6.77)	13.03
Strand 2012	8	249	0	128	-		•	9.05 (0.52, 157.98)	1.87
Dougados 1993	0	55	0	55				(Excluded)	0.00
Heyworth 2008	0	20	0	18				(Excluded)	0.00
Lundsgaard 2008	0	84	0	84				(Excluded)	0.00
Subtotal (I-squared	= 0.0%, 1	o = 0.39	7)			$\Diamond$		1.78 (1.10, 2.89)	65.53
Overall (I-squared :	= 0.0%, p	= 0.799	))			$\diamond$		1.78 (1.21, 2.63)	100.00
· ·								,	
NOTE: Weights are	from rand	dom effe	ects analys	s					
				0.01	0.1	1	15		
				Foyour		Descent			

**Fig. 5** Forest plot displaying the results of the meta-analyses comparing serious adverse events with intra-articular hyaluronic acid versus placebo in patients with OA: overall analysis and analyses on studies

with and without concomitant anti-OA medication allowed. CI confidence interval, OA osteoarthritis

the odds of AEs associated with IAHA, because most of the studies included in the analyses did not report all treatmentemergent AEs in both the intervention and control groups [73, 82, 89, 92]. Consequently, assessing the certainty of evidence using the GRADE approach, we found "low" to "moderate" certainty with all the outcomes evaluated, none of the results being associated with a "high" certainty of evidence.

For SOC comparisons, in our analysis, no statistically significant difference was found between IAHA injections and placebo for all categories, with the exception of infections and infestations. Overall, in the IAHA treatment group, there was significant lower odds of infections and infestations compared with the placebo group (OR = 0.61,

95% CI 0.40–0.93). This was also the case in the group of studies without concomitant pharmacological OA treatment (OR = 0.49, 95% CI 0.27–0.89). This has significantly been reported by a single study (Jorgensen et al. [65]), for which the author provided us with the full safety report and in which the main events reported were influenza, urinary tract infection, and pneumonia. This study counted for almost half (47.09%) of the weight of studies included in the overall analysis. A second study (with concomitant anti-OA medication) representing 49.84% of the overall studies weight also reported fewer infections in the IAHA group compared with placebo, but the difference in odds was not statistically significant (Jubb et al. [82]). A few in vitro studies suggest an anti-microbial effect of HA at levels of 1 mg/mL

# Intra-articular hyaluronic acid: Any adverse event

	n	Ν	n	Ν			%
Study	Active	Active	Placebo	Placeb		Odds Ratio (95% CI)	Weight
Without concomitant	OA treat	ment			1		
Baltzer 2009	51	135	30	107	+++-	1.56 (0.90, 2.69)	9.24
Brandt 2001	76	114	74	112		1.03 (0.59, 1.78)	9.11
Dixon 1988	3	30	0	33	<u> </u>	8.53 (0.42, 172.27)	0.38
Jorgensen 2010	78	165	77	170		1.08 (0.70, 1.66)	13.31
Puhl 1993	4	102	5	107		0.83 (0.22, 3.19)	1.84
Van der Weegen 20	15 30	99	25	97		1.25 (0.67, 2.34)	7.42
Subtotal (I-squared	= 0.0%, p	= 0.648	)		$\diamond$	1.19 (0.92, 1.54)	41.30
					l de la companya de l		
With concomitant OA	A treatmer	nt					
Altman 2009	157	293	169	295	-	0.86 (0.62, 1.19)	19.09
Dougados 1993	4	55	1	55		4.24 (0.46, 39.17)	0.69
Jubb 2003	187	208	168	200		1.70 (0.94, 3.06)	8.20
Kwon 2013	85	150	99	150		0.67 (0.42, 1.08)	11.78
Lundsgaard 2008	0	84	1	84		0.33 (0.01, 8.20)	0.33
Pham 2004	107	131	69	85		1.03 (0.51, 2.08)	6.10
Strand 2012	172	249	81	128		1.30 (0.83, 2.03)	12.50
Heyworth 2008	0	20	0	18	l.	(Excluded)	0.00
Subtotal (I-squared	= 40.1%,	p = 0.12	4)		$\diamond$	1.04 (0.77, 1.40)	58.70
Overall (I-squared =	17.7%, p	= 0.265	j)		6	1.09 (0.90, 1.31)	100.00
NOTE: Weights are	from rand	om effec	ts analysis	S			
					0.01 0.1 1 15	180	
					Favours intervention Does not favour	intervention	

**Fig. 6** Forest plot displaying the results of the meta-analyses comparing total adverse events with intra-articular hyaluronic acid versus placebo in patients with OA: overall analysis and analyses on studies

with and without concomitant anti-OA medication allowed. CI confidence interval, OA osteoarthritis

and over [97, 98]. These levels are unlikely to be achieved systemically following IA injection. However, if the results provided by the two RCTs demonstrating a lower rate of infections with IAHA versus placebo are not due to chance, they could suggest a systemic exposure of HA administered by IA injection. Further data are therefore needed to clarify this.

Given our findings overall, further investigation into the safety of IAHA is warranted. There are already a large number of studies that have assessed the efficacy and safety of IAHA in OA; the main issue is not therefore the lack of studies, but the lack of transparency in the reporting of IAHA safety data. Particularly, it would be interesting if the pharmaceutical companies could cooperate in future meta-analyses by giving access to full safety reports from the studies. Additionally, long-term safety studies are warranted as IAHA is often given as repeated courses of three to five injections; in fact, for most of the studies included in this systematic review, the follow-up durations varied from 8 to 26 weeks (Tables 1, 2). Multiple courses of IAHA are shown to be safe over 6–18 months from a postmarketing registry of one HA product (Supartz), with an overall AE rate of 0.008 (95% CI 0.001–0.055) [37]. The majority of people who reported an AE did so in the first injection series, of which 85% were injection site reactions. Conversely, an increased frequency of acute local reactions has been reported with multiple cycles of IAHA [99], while one review reports that the safety remains unchanged with



IAHA: Withdrawals due to adverse events

**Fig. 7** Forest plot displaying the results of the meta-analyses comparing withdrawals due to adverse events with IAHA versus placebo in patients with OA: overall analysis and analyses on studies with and

without concomitant anti-OA medication allowed. CI confidence interval, IAHA intra-articular hyaluronic acid, OA osteoarthritis

multiple courses of treatment [100]. Additionally, it has been reported that, in comparison with other pharmacological treatments for OA, IAHA appears to be associated with fewer systemic AEs than paracetamol or NSAIDs, but more local reactions, and a lower rate of withdrawals due to AEs [16, 17]. However, all this evidence deserves further investigation; transparency in the reporting of harms collected during clinical trials on IAHA will assist significantly in clarifying the safety profile of IAHA, for which the onus is on pharmaceutical companies developing HA for patients with OA.

#### 4.1 Strengths

Only RCTs versus placebo were included, and hence, the real effect is not underestimated. Many SOCs were investigated; not only "total AEs", "SAEs" or "skin AEs", as reported in many previous meta-analyses. To avoid double counting of AEs, for each SOC, we considered the number of patients



Harbord's test: p = 0.26

**Fig.8** Assessment of publication bias: funnel plot using data for the meta-analysis comparing total adverse events with intra-articular hyaluronic acid versus placebo in patients with OA. Harbord's test: p = 0.26. OA osteoarthritis, OR odds ratio

who experienced at least once any related AE. For total AEs (any AEs), we considered the number of patients who experienced, at least once, any AE during the study.

### 5 Limitations

Many studies identified that met the inclusion criteria did not provide AE data suitable for inclusion in the meta-analysis and the authors/sponsors did not provide us with the full safety data. As a consequence, subgroup analyses were not possible, which would have helped to explore any subgroup differences in the occurrence of SAEs with the studies included in the parallel post hoc meta-analysis (e.g. analyses by HA subtypes).

#### 6 Conclusions

Paucity in the reporting of AEs does not allow for a definitive conclusion regarding the safety profile of IAHA in OA. Based on the available data regarding studies without concomitant anti-OA medications, IAHA seems not to be associated with any safety issue in the management of OA; however, the certainty of this evidence was graded between "low" and "moderate". The SAEs found in some other metaanalyses as well as in our parallel post hoc analysis might be due to the allowance for concomitant use of oral NSAIDs during some trials or to other factors, but this deserves further investigation. Overall, further investigation on the safety of IAHA is warranted, in particular, greater contribution from pharmaceutical companies in providing full safety reports for future meta-analyses. Authors of manuscripts on future trials on IAHA are also encouraged to report harms collected during these trials in a transparent way. Further studies are also required to determine the long-term safety of IAHA.

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Outcomes	No. of participants	Certainty of the	Overall relative effect	Anticipated absolute effects	
	(studies), follow-up	evidence (GRADE)	) (95% CI)	Risk with placebo	Risk difference with IAHA
Gastrointestinal adverse events	1936 (11 RCTs)	⊕⊕⊕⊖ MODERATE	0.81 (0.52–1.27)	87 per 1000	15 fewer per 1000 (40 fewer to 21 more)
Cardiac disorders	2560 (13 RCTs)	⊕⊕⊖O LOW <sup>a,b</sup>	1.25 (0.36–4.41)	3 per 1000	1 more per 1000 (2 fewer to 11 more)
Vascular disorders	2560 (13 RCTs)	⊕⊕⊖O LOW <sup>a,b</sup>	1.70 (0.39–7.29)	2 per 1000	2 more per 1000 (1 fewer to 15 more)
Respiratory, thoracic and mediastinal disorders	2560 (13 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	1.21 (0.82–1.78)	42 per 1000	8 more per 1000 (7 fewer to 30 more)
Nervous system disorders	2089 (11 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	1.15 (0.77–1.70)	54 per 1000	8 more per 1000 (12 fewer to 35 more)
Skin and subcutaneous tissue disorders	1173 (8 RCTs)	⊕⊕⊖⊖ LOW <sup>a,b</sup>	1.71 (0.52–5.63)	18 per 1000	12 more per 1000 (8 fewer to 74 more)
Musculoskeletal and connective tissue disorders	2962 (13 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	0.99 (0.71–1.39)	101 per 1000	1 fewer per 1000 (27 fewer to 34 more)
Renal and urinary disorders	2560 (13 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	0.54 (0.21–1.41)	10 per 1000	4 fewer per 1000 (8 fewer to 4 more)
Infections and infestations	2019 (11 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	0.61 (0.40–0.93)	65 per 1000	24 fewer per 1000 (38 fewer to 4 fewer)
Hypersensitivity reaction	2560 (13 RCTs)	⊕⊕⊖⊖ LOW <sup>a,b</sup>	0.64 (0.05–7.94)	15 per 1000	5 fewer per 1000 (14 fewer to 94 more)
Severe adverse events	3232 (14 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	1.08 (0.50–2.31)	16 per 1000	1 more per 1000 (8 fewer to 20 more)
Serious adverse events	3956 (16 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	1.78 (1.21–2.63)	22 per 1000	17 more per 1000 (5 more to 34 more)
Withdrawals due to adverse events	3540 (15 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	1.62 (1.04–2.51)	26 per 1000	15 more per 1000 (1 more to 36 more)
Any adverse event	3476 (14 RCTs)	⊕⊕⊕⊖ MODERATE <sup>a</sup>	1.09 (0.90–1.31)	487 per 1000	22 more per 1000 (26 fewer to 67 more)

estimate is limited; the true effect may be substantially different from the estimate of the effect; Very low certainty: we have very little confidence in the effect estimate; the true effect is likely to be substantially different from the estimate of effect

The risk in the intervention group and its 95% CI are based on the assumed risk in the comparison group and the relative effect of the intervention and its 95% CI

CI confidence interval, OR odds ratio, RCT randomized controlled trial

<sup>a</sup>High risk of incomplete outcome data (attrition bias) and selective outcome reporting (reporting bias) found in many of the included studies. Full safety data were provided by authors of manuscripts for only 2 studies

<sup>b</sup>Wide CI because of low number of events

#### **Compliance with Ethical Standards**

All authors meet the International Committee of Medical Journal Editors (ICMJE) criteria for authorship for this manuscript, take responsibility for the integrity of the work as a whole, and have given final approval to the version to be published.

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