Is Local Viscosupplementation Injection Clinically Superior to Other Therapies in the Treatment of Osteoarthritis of the Knee: A Systematic Review of Overlapping Meta-analyses

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Purpose: To conduct a systematic review of overlapping meta-analyses comparing treatment of knee osteoarthritis (OA) with intra-articular viscosupplementation (intra-articular hyaluronic acid [IA-HA]) versus oral nonsteroidal antiinflammatory drugs (NSAIDs), intra-articular corticosteroids (IA-corticosteroids), intra-articular platelet-rich plasma (IA-PRP), or intra-articular placebo (IA-placebo) to determine which meta-analyses provide the best current evidence and identify potential causes of discordance. **Methods:** Literature searches were performed for meta-analyses examining use of IA-HA versus NSAIDs, IA-corticosteroids, IA-PRP, or IA-placebo. Clinical data were extracted, and meta-analysis quality was assessed. The Jadad algorithm was applied to determine which meta-analyses provided the highest level of evidence. Results: Fourteen meta-analyses met the eligibility criteria and ranged in quality from Level I to IV evidence. In studies reporting patient numbers, there were a total of 20,049 patients: 13,698 receiving IA-HA, 355 receiving NSAIDs, 294 receiving IA-corticosteroids, and 5,702 receiving IA-placebo. Ten studies examined the effects of IA-HA versus IA-placebo; of these, 5 found that IA-HA improved pain and 4 found that IA-HA improved function. No clinically relevant differences in the efficacy of IA-HA versus NSAIDs regarding pain and function were found. Regarding IA-HA versus IA-PRP, IA-HA improved knee function at 2 and 6 months after injection but the effects were less robust than those of IA-PRP. Regarding IA-HA versus IA-corticosteroids, the positive effects of IA-HA were greater at 5 to 13 weeks and persisted for up to 26 weeks. After application of the Jadad algorithm, 2 concordant high-quality meta-analyses were selected and both showed that IA-HA provided clinically relevant improvements in pain and function compared with IA-placebo. Conclusions: This systematic review of overlapping meta-analyses comparing IA-HA with other nonoperative treatment modalities for knee OA shows that the current highest level of evidence suggests that IA-HA is a viable option for knee OA. Its use results in improvements in knee pain and function that can persist for up to 26 weeks. IA-HA has a good safety profile, and its use should be considered in patients with early knee OA. Level of Evidence: Level IV, systematic review of Level I to IV studies.

K nee pain due to osteoarthritis (OA) is one of the most common complaints in patients presenting to orthopaedic clinics, resulting in significant societal costs including cost of treatment and lost time from work or

activities.^{1,2} Several nonoperative and operative treatment options exist to mitigate this pain and the resulting limitations in function occurring in patients with arthritis. The goal of nonoperative treatment modalities is to minimize

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The authors report the following potential conflict of interest or source of funding: B.R.B. receives support from Arthrex, Ossur, Linvatec, and Smith \mathcal{P} Nephew. B.J.C. receives support from Arthrex, DJ Orthopaedics, Johnson \mathcal{P} Johnson, Regentis, Zimmer, and Smith \mathcal{P} Nephew. N.N.V. receives support from Minivasive, Smith \mathcal{P} Nephew, Arthrosurface, Omeros, Arthrex, Athletico, ConMed Linvatec, Miomed, and Mitek.

Received December 10, 2014; accepted March 19, 2015.

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^{© 2015} by the Arthroscopy Association of North America 0749-8063/141037/\$36.00 http://dx.doi.org/10.1016/j.arthro.2015.03.030

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pain and restore function in a noninvasive manner while prolonging the need for a total knee arthroplasty (TKA). These options include intra-articular viscosupplementation (intra-articular hyaluronic acid [IA-HA]), intra-articular corticosteroids (IA-corticosteroids), oral nonsteroidal antiinflammatory drugs (NSAIDs), and intra-articular platelet-rich plasma (IA-PRP).

Viscosupplementation is the injection of an intraarticular compound made of high-molecular-weight fluid containing hylan products (derivative of hyaluronan) that essentially functions as a viscoelastic glycosaminoglycan. Hyaluronic acid (HA) is naturally present in joint fluid and serves multiple purposes including shock absorption, joint lubrication, and energy dissipation; in addition, it coats the articular cartilage surfaces of the femur, tibia, and patella to protect them.³

The desire to delay the treatment of knee OA with TKA lies in the desire to reduce the possibility of the need for early revision TKA. Although the failure rate varies on an individual basis, it is generally accepted that the revision rate for knee arthroplasty is slightly less than 1% per year with a 10-year survivorship rate of approximately 95% and a 20-year survivorship rate of approximately 85%.⁴⁻⁸ Recent evidence has shown that approximately 4 million persons in the United States are living with a TKA and that over half of the adults in the United States diagnosed with knee OA will eventually undergo TKA.⁹

Despite the plethora of studies examining the array of less invasive treatment options that exist for knee OA prior to performing a TKA, there has been no definitive consensus as to which treatments are the most effective at improving pain and function.^{10,11} Arrich et al.¹⁰ performed a meta-analysis to determine if IA-HA improved pain or function in patients with knee OA and found that it did improve activity-related knee pain. Conversely, Bannuru et al.¹¹ conducted a meta-analysis comparing IA-HA with oral anti-inflammatory medications, and although both treatments showed improvements in function and stiffness, there were no differences between the groups.

Therefore the purpose of this study was to conduct a systematic review of overlapping meta-analyses comparing treatment of knee OA with IA-HA versus oral NSAIDs, IA-corticosteroids, IA-PRP, or intra-articular placebo (IA-placebo) to determine which meta-analyses provide the best current evidence and identify potential causes of discordance. The main objectives of this study were (1) to conduct a systematic review of meta-analyses comparing the aforementioned treatment options for knee OA, (2) to provide an analytical framework for interpreting the presently discordant best available evidence to develop treatment recommendations, and (3) to identify gaps in the literature that require continued investigation. We hypothesized that intra-articular injections of HA would provide significant improvement in

pain and function with minimal side effects compared with IA-corticosteroids, IA-PRP, IA-placebo, or oral antiinflammatory medications.

Methods

A systematic review of the literature was performed using the PubMed database, CINAHL (Cumulative Index to Nursing and Allied Health Literature) Complete database, Cochrane Database of Systematic Reviews, Scopus database, and Embase database. The following search terms were used: meta-analysis AND hyaluronic acid AND (knee [arthritis OR osteoarthritis]) AND (corticosteroid OR NSAID OR placebo OR [platelet rich plasma OR PRP]). The search was performed on August 24, 2014, and was limited to articles written in English. Broad search query terms were used to include all possibly applicable studies. All reviewed articles were then manually cross-referenced to ensure that all potential studies were included.

The abstracts that resulted from these searches were reviewed by 2 of the authors (K.A.C. and R.M.). The inclusion criteria were meta-analyses that compared the use of IA-HA in knee OA with the use of IAplacebo, IA-PRP, IA-corticosteroids, or oral NSAIDs. Cadaveric, animal, and biomechanical studies were excluded. The exclusion criteria included narrative reviews, reviews without an organized and reported search algorithm, reviews that did not directly compare IA-HA versus another treatment modality, studies without clinical outcome data, and non-Englishlanguage studies. Systematic reviews that did not pool data or perform a meta-analysis were also excluded. Full-text articles were then obtained for those studies that met both the inclusion and exclusion criteria. The references for each of these citations were manually screened to ensure that no studies were missed. The tables of contents for the past 2 years of Arthroscopy, The Journal of Bone and Joint Surgery, The American Journal of Sports Medicine, Clinical Orthopaedics and Related Research, Osteoarthritis and Cartilage, and The New England Journal of Medicine were manually searched for any additional studies that were not identified in our prior search. A PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) diagram shows our study selection algorithm (Fig 1).

Data were extracted from the studies that met the inclusion criteria and included information about levels of evidence included in the studies, length of follow-up, duration of symptomatic relief, adverse events, knee function, knee pain outcomes, and pooled effect size. Standardized outcome scores that were collected included Lequesne scores, visual analog scale (VAS) pain scores, and Western Ontario and McMaster Universities Osteoarthritis Index pain subscores. Data specific to the methodology of the included meta-analyses

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Viscosupplementation Meta-Analysis

Fig 1. PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) diagram showing the results of application of the study algorithm to the number of studies included, with the number of studies removed after application of each exclusion criterion. (NSAID, nonsteroidal antiinflammatory drug; PRP, platelet-rich plasma.)



were extracted and included the rationale for repeating the systematic review, the databases that were used for the review, a comparison of the number of "possible" previous systematic reviews cited versus the number that were actually cited in the study, and the conclusions of the review regarding whether IA-HA was more clinically effective than the treatment modality with which it was compared in terms of pain relief and adverse events.

Meta-analysis quality was scored using the Quality of Reporting of Meta-analyses (QUOROM) system.¹² This system provides a method for evaluating meta-analyses based on the quality of their reporting and methodology in 18 categories. Each meta-analysis was awarded a point in each category if it met over half of the criteria given in that category, for a total of 18 points possible. Meta-analysis quality was also graded using the Oxman-Guyatt quality-appraisal tool.¹³ The modified Coleman

Methodology Score¹⁴ was extracted from individual studies when available. In addition, when known biases within the literature were reported by individual trials, these were recorded.

The Jadad decision algorithm¹⁵ was used to guide interpretation of discordant reviews. Sources of discordance among meta-analyses as described by Jadad et al.¹⁵ include differences in the clinical question, inclusion/exclusion criteria, data pooling, data extraction, quality assessment, and statistical analysis. Scoring was performed based on the assessment of randomization, randomization methodology, double blinding, withdrawals or dropouts from the study, and allocation concealment. This algorithm was independently applied by 3 of the authors, and their results were compared to determine which of the included systematic reviews provided the best current evidence to make recommendations (K.A.C., B.M.S., R.M.). All statistical

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analyses were performed with the use of Microsoft Excel X (Microsoft, Redmond, WA).

Results

The initial search yielded 105 abstracts, and after application of the study selection algorithm, 14 studies fulfilled our inclusion and exclusion criteria and were included (Fig 1). These studies were published between 2003 and 2014, with all 14 performing a meta-analysis. Industry funding for the studies was provided to 4 of the 14 included meta-analyses,¹⁶⁻¹⁹ which introduces the possibility of a conflict of interest.

Of the studies, 7 included Level I evidence,^{11,17,18,20-23} 4 included Level I and Level II evidence,^{10,19,24,25} 1 included Level II and Level III evidence,²⁶ and 2 included Level I and Level IV evidence.^{16,27} The number of included patients ranged from 606 patients²³ to 12,667 patients,²⁵ with mean follow-up periods ranging from 3 weeks²¹ to 135.2 weeks.¹⁶ In studies that reported the number of patients in each group, there were a total of 20,049 patients: 13,698 receiving IA-HA, 355 receiving NSAIDs, 294 receiving IA-corticosteroids, and 5,702 receiving IA-placebo.

Authors' Assessment of Prior Systematic Review Literature

The majority of the included studies only cited a few of the available pre-existing meta-analyses or systematic reviews (Table 1), with only 7 of 14 studies citing more than 50% of prior systematic reviews or metaanalyses.^{10,11,19,21-23,25} None of the studies cited all of the previous systematic reviews that were available at the time of publication. Eleven of the 14 studies provided a rationale for repeating the systematic review (Tables 1 and 2), with several of them highlighting the discordant findings of prior meta-analyses as the rationale for repeating the study.^{17,22} The remaining studies cited either differing methodologies,^{19,22} the inclusion of other outcome variables,^{10,18,20} or a comparison with other treatments as their rationale.^{11,21,23,27} Appendix Table 1 (available at www.arthroscopyjournal.org) provides a list of the primary studies used in each metaanalysis.

Outcome Measures

The included studies were heterogeneous in both the standardized and nonstandardized patient outcome measures that were reported (Appendix Table 2, available at www.arthroscopyjournal.org). There was a high level of variance seen in the mean differences in VAS scores (on a 100-mm VAS) for IA-HA versus IA-placebo. The studies by Arrich et al.¹⁰ and Modawal et al.,²⁰ in which the mean differences ranged from -3.8 to 18.1, highlight these differences. In terms of a weighted mean difference between the efficacy of IA-HA and intra-articular saline solution, a value of just

10.20 using the VAS was found at 3 months' follow-up,¹⁸ whereas standardized mean difference (SMD) values representing only a small (SMD <0.5) effect on knee pain were found at follow-up.^{19,24} For reference, the SMD values of 0.2, 0.5, 0.8, and 1.0 are defined as small, medium, large, and very large, respectively.²⁸

When IA-HA was compared with IA-placebo, the patient-reported mean percent improvement in pain from baseline at the 5- to 13-week post-injection time point ranged from 28% to 54% whereas the mean percent improvement in function ranged from 9% to 32%.²² In terms of the effect size for IA-HA versus NSAIDs, no clinically relevant differences were found between the 2 treatments.^{11,16} IA-PRP was found to have a greater pooled effect size regarding knee function (when pooled using a random-effects model) compared with IA-HA at both 2 months and 6 months after injection. This effect was maintained for up to 1 year.²⁷

The included studies were also heterogeneous with respect to their method of analysis of the response to treatment with IA-HA. Some studies reported on pain outcome scores and knee function, whereas others reported on knee range of motion, activity-related knee pain, pooled effect size, VAS pain score, Lequesne score, Western Ontario and McMaster Universities Osteoarthritis Index score, or adverse reactions (Appendix Table 3, available at www.arthroscopyjournal.org).

Search Methodology

Although 13 of the 14 included studies searched PubMed/Medline, there was heterogeneity in the other databases that were used. These included Embase, the Cochrane Database of Systematic Reviews, and the CINAHL among others. One study did not identify the databases used in the search.¹⁷ Ten of the included studies used the Embase or Cochrane Database of Systematic Reviews (or both) to find articles. Three studies used the CINAHL database, 10,21,23 and all studies that reported their search strategy searched at least 3 databases (Table 2). The total number of unique primary studies cited by the included systematic reviews was 107. The number of primary studies varied widely from 5 in those reviews performed in 2006 and 2014^{11,17} to 89 in a study published in 2012,²⁵ with a median of 18 primary studies cited (Table 2 and Appendix Table 1 [available at www.arthroscopyjournal.org]).

Study Results

IA-HA Versus IA-Placebo. Of the 10 studies that examined the effects of IA-HA versus IA-placebo, 5 found that IA-HA resulted in improvements in pain and 4 found that it resulted in improvements in function. However, 3 meta-analyses found no difference between IA-HA and IA-placebo in terms of pain, and

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Table 1. Number of Prior Systematic Reviews or Meta-analyses Actually Cited as Compared With Maximum Number That CouldPossibly Have Been Cited, in Addition to Authors' Rationale for Repeating Systematic Review

Authors	Date of Publication (Month/Day/Year)	Date of Last Literature Search (Month/Day/Year)	No. of Systematic Reviews or Meta-analyses Possible to Cite	No. of Systematic Reviews or Meta-analyses Cited	Rationale for Repeating Meta-analysis as Abstracted From Article
Espallargues	1/—/2003	—/—/1999	0	0	NA
and Pons ¹⁶ Lo et al. ²⁴	12/17/2003	2/—/2003	1	0	NA
Wang et al. ²⁰	3/—/2004	12/—/2001	0	0	NA
Arrich et al.**	4/12/2005	4/—/2004	3	2	"In contrast to 2 previous meta-analyses on this subject, we used a different approach to data synthesis and interpretation: instead of analyzing a composite effect size over time, we allocated trial data, when possible, to 3 outcome groups that we assumed would be relevant for patients with osteoarthritis. We specifically looked at pain at rest, pain during exercise and joint function as distinct outcomes, measured repeatedly over time. In addition, we assessed adverse events and the impact of both trial quality and molecular mass of the product. This analysis allows us to provide important additional insight into the effects of intra-articular administration of hyaluronic acid for the treatment of osteoarthritis of the knee."
Modawal et al. ²⁰	9/—/2005	8/—/2004	3	1	"We provide here a stringent test of the efficacy of viscosupplementation for relieving knee pain from osteoarthritis with a meta-analysis that includes only data from randomized, double- blinded, controlled trials of hyaluronic acid that measured pain using a visual analogue scale (VAS), the most widely accepted method for pain evaluation."
Strand et al. ¹⁷	<i>—/—/2006</i>	NA	5	2	"Divergent interpretations from 3 recent meta- analyses have added to this controversy To supplement evidence provided by recent meta- analyses of IA-HA treatment, an integrated analysis of five RCTs examining a single IA-HA product is presented This provides a comparison that avoids some limitations inherent to meta-analyses, because it circumvents the need for any type of data transformation "
Reichenbach et al. ²¹	12/15/2007	11/—/2006	6	4	"All of these studies compared hyaluronic acid and hylan with a sham intervention, but only one study included trials comparing hylan with hyaluronic acids directly heterogeneity of the studies limited conclusions did not pool results of included trials The safety of hylan compared with conventional hyaluronic acids was rarely addressed but sample sizes of included trials precluded any definitive conclusions Previous claims that hylan has greater benefits compared with conventional preparations of hyaluronic acids were mainly based on implicit indirect comparisons from placebo-controlled trials."
Bellamy et al. ²²	—/—/2009 (update—original 4/19/2006)	1/—/2006	6	5	"These publications employ different methodologies and have shown conflicting results but they recommended further work on the effect of multiple courses of hylan Given this diversity of opinion there is, therefore, a rational basis for performing a Cochrane review of viscosupplementation in knee OA."

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Table 1. Continued

			No. of	No. of	
	Date of	Date of Last	Systematic Reviews or	Systematic Reviews or	Rationale for Repeating
	Publication	Literature Search	Meta-analyses	Meta-analyses	Meta-analysis as
Authors	(Month/Day/Year)	(Month/Day/Year)	Possible to Cite	Cited	Abstracted From Article
Bannuru et al. ²³ (2009)	12/15/2009	2/—/2009	8	5	"However, the conclusions of meta-analyses were also inconsistent In the face of this controversy, we aimed to reexamine the clinical usefulness of HA products from the perspective of their relative efficacy when compared with intraarticular corticosteroids, a widely used intervention with which clinical rheumatologists have considerable familiarity."
Rutjes et al. ²⁵	6/12/2012	1/31/2012	9	7	"Several trials have since been published [since previous reviews]. In addition, we were aware of unpublished trials, which were never included in any meta-analysis to date. Therefore, we did a comprehensive, up-to-date systematic review to determine whether viscosupplementation is clinically effective and safe to treat symptomatic knee OA."
Colen et al. ¹⁸	8/1/2012	6/27/2011	9	4	"In this systematic review we will compare the efficacy of intra-articularly administered HA with intra- articularly administered placebo in randomized controlled trials (RCTs) using the visual analog scale (VAS) for pain as a primary outcome measurement at 3-months follow-up. Using this approach we make some recommendations concerning the efficacy of HA compared with the effects of placebo and discuss the differences in efficacy between the different HA products and the differences between the different HA products and placebo."
Miller and Block ¹⁹	9/1/2013	6/—/2013	11	8	"In contrast, we only included data from full-text manuscripts published in peer-reviewed journals. Lastly, Rutjes et al analyzed all safety data using an odds ratio, a statistic that excludes zero total event trials. Considering that 30 of 38 SAE treatment effects in the current meta- analysis reported zero total events, the odds ratio is arguably an inappropriate statistic for this type of analysis since most data are disregarded."
Chang et al. ²⁷	3/—/2014	9/—/2013	12	4	"However, to our knowledge, no meta-analytic research has quantified the effectiveness of PRP treatment and analyzed the factors that modify the outcomes. Therefore, we undertook a systematic review and meta-analysis to investigate the clinical results in patients with knee chondral degenerative lesions, with regard to functional changes, compared with the pretreatment condition, after PRP injections, placebo controls and HA administration "
Bannuru et al. ¹¹ (2014)	4/—/2014	4/—/2013	13	9	"Several meta-analyses have examined the effects of IA-HA in the treatment of knee OA compared with placebo and with intra-articularly injected corticosteroids and found inconclusive results Although NSAIDs are among the most efficacious and widely used treatments for knee OA, no meta- analysis has been performed to assess these medications against IA-HA, which is considered to have a more favorable safety profile. The objectives of this study were to systematically evaluate the relative efficacy of IA-HA for symptomatic knee OA in comparison with NSAIDs "

HA, hyaluronic acid; IA-HA, intra-articular hyaluronic acid; NA, not available; NSAIDs, nonsteroidal anti-inflammatory drugs; OA, osteoarthritis; PRP, platelet-rick plasma; RCTs, randomized controlled trials; SAE, serious adverse event.

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Table 2. Search Methodology Used by Included Studies

			Dat	tabase				
Authors	PubMed/ Medline	Embase	Cochrane Library	CINAHL	Science Citation Index	Other	No. of Primary Studies	Primary Studies Included Only RCTs
Espallargues and Pons ¹⁶	+	+	+	_	—	+	14	-
Lo et al. ²⁴	+	—	+	—	_	+	22	+
Wang et al. ²⁶	+	+	+	-	-	+	20	+
Arrich et al. ¹⁰	+	+	+	+	_	+	22	+
Modawal et al. ²⁰	+	-	_	-	_	_	9 (11 comparable cohorts)	+
Strand et al. ¹⁷	NR	NR	NR	NR	NR	NR	5	+
Reichenbach et al. ²¹	+	+	_	+	+	+	13 (15 comparable cohorts)	+
Bellamy et al. ²²	+	+	+	-	-	+	76	-
Bannuru et al. ²³ (2009)	+	+	+	+	_	+	7	+
Rutjes et al. ²⁵	+	+	+	-	+	+	89	-
Colen et al. ¹⁸	+	+	+	-	-	_	74	+
Miller and Block ¹⁹	+	+	_	-	_	_	29	+
Chang et al. ²⁷	+	_	+	-	-	+	16	-
Bannuru et al. ¹¹ (2014)	+	+	+	-	_	+	5	+

CINAHL, Cumulative Index to Nursing and Allied Health Literature; Embase, Excerpta Medica Database; Medline, Medical Literature Analysis and Retrieval System Online; RCTs, randomized controlled trials; NR, not recorded.

4 studies found no difference in function. The remaining studies showed no clinically relevant differences in either pain or function.^{10,16-20,22,24-26}

IA-HA Versus Oral NSAIDs. No clinically relevant differences in the efficacy of IA-HA versus oral NSAIDs on knee pain and function were found in the 3 studies that examined it. However, IA-HA was found to have a slightly more favorable adverse reaction profile than NSAIDs because of the risk of gastrointestinal side effects posed by NSAIDs.^{11,16,22} Although both IA-HA and IA-PRP led to improvements in knee function at 2 and 6 months after injection, the positive effects of IA-HA were less robust than those of IA-PRP and there were no differences in adverse reactions.²⁷

IA-HA Versus IA-Corticosteroids. IA-corticosteroids provided better pain relief during the first 4 weeks after injection, but the positive effects of IA-HA were greatest at the 5- to 13-week post-injection time point, and this relief persisted for up to 26 weeks in 2 studies.^{22,23} No definitive conclusions could be drawn about the best HA product in the studies that compared the different formulations of HA products.^{16,18,21,22} In 1 study comparing IA-HA versus intra-articular hylan,²¹ the authors discouraged the use of intra-articular hylan because of the increased risk of adverse reactions and their finding of no clinically relevant evidence to support its use.

Study Quality and Validity

The QUOROM scores were assessed for each of the studies and ranged from 11^{16} to 17, 22,27 with a median of 15.5 (with the maximum possible score being 18). The Oxman-Guyatt scores ranged from 3^{16} to $7^{22,27}$ on a scale from 1 to 7, with a median score of 5. As a reference, Oxman-Guyatt scores lower than 3 are

generally considered to indicate that the study in question has "major flaws."¹³

Heterogeneity Assessment

Several methods were used to assess study heterogeneity, and 12 of the 14 included studies performed a statistical heterogeneity analysis.^{10,11,18-27} Several performed subgroup analyses assessing parameters such as pain outcomes, physical function, pain by VAS score, pain at rest, pain with activities, and major adverse effects (Appendix Table 2, available at www. arthroscopyjournal.org). Given the heterogeneity in the variables examined, treatment time point assessed, and overall findings of the subgroup analyses that were performed, it was found that the 2 highest-quality studies^{22,27} provided the best available evidence about the use of IA-HA in knee arthritis.

Application of Jadad Decision Algorithm

The Jadad decision algorithm was applied to determine which of the 14 included meta-analyses provided the best available current evidence for treatment recommendations in patients with knee OA. The 3 authors applying the Jadad algorithm independently selected the same route through the Jadad decision algorithm. Given that (1) all of the meta-analyses did not address the same study question, (2) our reviews did not include the same primary trials (Table 2 and Appendix Table 1 [available at www.arthroscopyjournal.org]), and (3) our reviews did not have the same selection criteria, the Jadad algorithm suggests that the highest-quality review can be selected based on the publication characteristics of the primary trials, the methodology of the primary trials, the language restrictions, and whether an analysis of data on individual patients was included in the study. The last 2 criteria do

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not apply to this study. With respect to publication status, several newer meta-analyses included multiple newly available trials, which may explain some of the discordance in the results and conclusions that were drawn. Regarding the methodology of primary trials, those reviews that included only Level I evidence included trials of superior methodology. Use of the aforementioned criteria facilitated the selection of 2 high-quality meta-analyses with results that represent the best available current evidence.^{22,27} These studies were not industry sponsored and concluded that IA-HA leads to improvement in knee pain and function in patients with knee OA versus IA-placebo. The positive effects of IA-HA versus IA-corticosteroids were greatest at 5 to 13 weeks after injection, and this effect persisted for up to 26 weeks after injection. No clinically relevant differences in the efficacy of IA-HA versus oral NSAIDs on knee pain and function were found, but consideration should be given for IA-HA use in patients with knee OA who are unable to tolerate NSAIDs. Although IA-HA leads to improved knee function at 2 and 6 months' follow-up, IA-PRP leads to greater improvements. No definitive conclusions could be drawn about the best HA product in the studies that compared the different formulations of HA products. Overall, the best available evidence in the literature supports the use of the HA class of products in the treatment of knee OA.

Discussion

This systematic review of overlapping meta-analyses found that intra-articular viscosupplementation is a safe and viable treatment option for knee OA with effects that can last up to 26 weeks. Given the high prevalence of knee OA, there have been multiple clinical trials, systematic reviews, and meta-analyses that have attempted to determine the best nonoperative treatment for this condition.⁹ However, a clear gold standard has not been identified. Therefore the main purposes of this systematic review of overlapping metaanalyses were to determine the source of discordance between the various meta-analyses and to determine which studies provided the best available current evidence on nonoperative treatment of knee OA. A critical inspection and assessment of the quality of the 14 included meta-analyses using the QUOROM and Oxman-Guyatt guidelines were undertaken to explore the best nonoperative treatment for knee OA. The included meta-analyses used studies of varied levels of evidence including 7 studies with Level I evidence,^{11,17,18,20-23} 4 with Level I and Level II evidence,^{10,19,24,25} 1 with Level II and Level III evidence,²⁶ and 2 with Level I and Level IV evidence.^{16,27} On the basis of the best available current evidence, the hypothesis that IA-HA provides significant improvement in pain and function with a minimal side-effect profile in the treatment of knee OA was confirmed.

With the changing health care environment, more emphasis has been placed on evidence-based medicine to determine the efficacy and cost-effectiveness of treatments to direct clinical practice guidelines. As such, it is important to identify the best evidence surrounding a particular treatment because there are a multitude of low-level studies with varying results on many different treatment modalities. The topic of nonoperative treatment for knee OA is extremely important because the number of patients with knee OA continues to grow.⁹ To minimize knee pain and functional limitations while delaying knee replacement, several nonsurgical options have been implemented, which include IA-HA, IA-corticosteroid injections, oral NSAIDs, and IA-PRP. Although each treatment has proved to relieve pain and improve function on its own, the question remains as to which treatment is superior in eliminating pain and improving function while providing a favorable side-effect profile. In this study we have determined that the highest level of evidence currently available supports the use of intra-articular viscosupplementation for the treatment of patients with knee OA.

The available evidence shows that IA-HA provides small but clinically relevant improvements in knee pain and function when compared with IA-placebo.^{10,16-} ^{20,22,24-26} Furthermore, although there were no major clinical differences in the efficacy of IA-HA versus oral NSAIDs in relieving knee pain and restoring function, the fact that IA-HA has a more favorable side-effect profile than oral NSAIDs makes IA-HA a good option for patients unable to tolerate oral NSAIDs.^{11,16,22} Both IA-HA and IA-corticosteroids were effective in controlling pain, with steroids providing better short-term relief and HA providing more long-term pain relief starting in the 5- to 13-week post-injection period and lasting for up to 26 weeks.^{22,23} Although the effects of IA-PRP were greater than those of IA-HA in terms of knee function at 2 and 6 months after injection,²⁷ the fact that platelet-rich plasma (PRP) is not currently reimbursed by insurance companies limits its availability. Treatment with PRP is cost prohibitive for most patients, and in light of the good outcomes achieved with IA-HA in terms of pain and function, IA-HA may be a better treatment option. Furthermore, more highquality randomized, double-blinded studies are needed to compare the effects of PRP versus other treatments before it can be fully endorsed as a viable universal option for patients with knee OA.

The recommendations from the recent American Academy of Orthopaedic Surgeons (AAOS) clinical practice guideline on the non-arthroplasty treatment of knee OA²⁹ highlights the need for more robust Level I evidence studies on all the aforementioned treatment modalities. Of the treatments investigated in this systematic review, oral NSAIDs were the only treatment modality recommended with a "strong" strength of

recommendation. The AAOS did not recommend the use of HA for patients with symptomatic OA of the knee and graded the strength of this recommendation as strong. The rationale for this rating was based on the fact that although statistically significant outcomes were seen in some studies, those outcomes were not clinically significant, based on a lack of minimum clinically important improvement-defined as the smallest clinical change that would be important to a patient—while also accounting for the fact that there are some statistically significant treatment-related improvements that may be too small to be clinically relevant, despite the fact that they are statistically significant. Similarly, the AAOS was unable to recommend for or against the use of IAcorticosteroids or IA-PRP and rated this recommendation as "inconclusive."²⁹ Interestingly, Bannuru et al.³⁰ highlighted that the AAOS clinical practice guideline's use of the minimum clinically important improvement metric may have contained some flaws, so despite the fact that IA-HA was not recommended based on this metric, the best available current evidence suggests that patients may still obtain some benefit from IA-HA use for knee OA. This discrepancy again highlights the need for more high-quality Level I studies exploring the different nonoperative treatment modalities for managing patients with knee OA, as well as the need for more studies on the use of the minimum clinically important improvement metric for these modalities.

The strengths of this study lie in the use of validated quality-assessment tools^{12,13,15} to critically appraise the studies included in our review. The use of these tools combined with their application in independent quality assessment by 3 authors with consensus agreement adds support to our main findings that intra-articular viscosupplementation is a viable treatment option for patients with knee OA.

Limitations

Similar to many of the studies that have been included in this review, there are some inherent limitations to our study. One of the major limitations is the fact that the quality of this systematic review is limited by the quality of the studies that were included in prior studies. As such, despite the fact that most of the included studies used data from Level I and Level II studies, the inclusion of Level IV evidence by 2 metaanalyses makes this a Level IV study by default. However, the findings should not be discredited because the conclusions of this study were based on 2 Level I studies.^{22,27}

Another limitation lies in the presence of heterogeneity in terms of the number of patients included in the studies, the type of interventions compared, the outcome measures collected, the type of subgroup analysis performed, and the use of the pooled effect size. In addition, the relatively short follow-up periods that were reported in some of the studies are a limiting factor, and little is known about the long-term effects of the described interventions on patients with knee OA. In this study the mean follow-up periods ranged from 3 weeks²¹ to 135.2 weeks,¹⁶ but most of the available outcome data regarding the impact of IA-HA on knee pain and function were only available for patients in the 2- to 3-month time period, with even fewer data available for 1 year. This lack of long-term data highlights the fact that more high-quality studies are needed to definitively determine how effective viscosupplementation is for patients with knee OA in the long-term.

Conclusions

According to this systematic review of overlapping meta-analyses comparing IA-HA with other nonoperative treatment modalities for knee OA, the current highest level of evidence suggests that IA-HA is a viable option for patients with knee OA. Its use results in improvements in knee pain and function that can persist for up to 26 weeks in comparison with other treatment modalities. IA-HA has been shown to have a good safety profile, and its use should be considered in patients with early knee OA.

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patients with knee osteoarthritis: Systematic review and meta-analysis of randomized, saline-controlled trials. *Clin Med Insights Arthritis Musculoskelet Disord* 2013;6:57-63.

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Appendix Table	1. Primary	Studies	Included	in	Meta-analysis	
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Primary Study	Espallargues and Pons ¹⁶	Lo et al. ²⁴	Wang et al. ²⁶	Arrich et al. ¹⁰	Modawal et al. ²⁰	Strand et al. ¹⁷	Reichenbach et al. ²¹	Bellamy et al. ²²	Bannuru et al. ²³ (2009)	Rutjes et al. ²⁵	Colen et al. ¹⁸	Miller and Block ¹⁹	Chang et al. ²⁷	Bannuru et al. ¹¹ (2014)
Adams, 1995	+	_	+	_	_	_	_	+	_	_	+	_	_	+
Altman, 1998	_	+	_	+	+	_	_	+	_	+	+	+	_	+
Altman, 2004	_	_	_	_	_	_	_	+	_	+	+	—	_	_
Altman, 2009	_	_	_	_	_	_	_	_	_	+	+	+	_	_
Anika, 2000	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Anika, 2001	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Ardic, 2001	_	_	_	_	_	_	_	+	_	+	_	_	_	_
Atamaz, 2004	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Atamaz, 2005	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Atamaz, 2006	_	_	_	_	_	_	+	_	_	_	+	_	_	_
Atay, 2008	_	_	_	_	_	_	_	_	_	+	+	_	_	_
Auerbach, 2002	_	_	_	_	_	_	_	+	_	_	+	_	_	_
Bach, 1997	+	_	_	_	_	_	_	_	_	_	_	_	_	_
Baltzer, 2009	_	_	_	_	_	_	_	_	_	+	+	_	_	_
Baraf, 2009	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Bayramoglu 2003	_	_	_	_	_	_	+	+	_	+	+	_	_	_
Beks 1997	+	_	_	_	_	_	_	_	_	_	_	_	_	_
Bellamy 2005	_	_	_	_	_	_	_	_	_	_	+	_	_	_
Blanco 2008	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Bragantini 1987	_	_	+	+	_	_	_	+	_	+	+	+	_	_
Brandt 2001	_	+	+	+	_	_	_	+	_	_	+	+	_	_
Brown 2003	_	_	_	_	_	_	_	- -	_	_	_	_	_	_
Bunyaratavei 2001	_	_	_	1	_	_	_	- -	_	1	_	-	_	_
Butup 2002	_	_	_	T	_	_	_		_	T 1	_	-	_	_
Caborn 2003			_							т _		_		
Caborn 2004	_		_	_		_	_	+	_	_	_	_	_	_
Caragual 2001		_						T	T	_	Ŧ		_	
Carrabba 1005	_		_	_	_	_	—	_	_	+	_	_	_	_
Corra 2012		Ŧ	Ŧ	Ŧ				T		Ŧ	Ŧ	Ŧ	1	
Cerza, 2012 Chavaliar, 2010	—	—	—	_	—	_	—	_	—	_	_	—	+	—
Cheve 2000										Ŧ	+			
	—	—	—	_	—	_	—	—	—	_	+	—	—	—
Cobap 1004	—	_	_	_	—	_	—	_	—	+	—	—	—	—
Conversion 2000	—	+	+	_	—	_	—	+	—	+	_	—	—	—
Conrozier, 2009	—	_	_	_		_	—	_	—	_	+		_	—
	—	+	+	+		_	—	+	—	+	+		_	—
Creatiler, 1994	—	+	+	_		_	—	+	—	+	+	_	_	—
	—	_	_	_		_	—	+	—	_	_	+	_	—
Cubukcu, 2005	_		_	_	_	_	_	_	_	+	_	_	_	_
Daniberg, 1994	—	+	_	+	_	_	—	_	—	_	+	_	_	—
Day, 2001	_	_	—		—		_	+	_				_	_
Day, 2004	—	_	_	+	_	+	—	+	_	+	+	+	_	—
DeCaria, 2012	—	_	_	_	_	_	—	_	_	_	_	+	_	—
Dickson, 1998	+	—	+	_	_	_	_	-	_	_	_	_	—	—
Dickson, 2001	—	_	_	_	_	_	_	+	_	+	_	_	_	+
Diracoglu, 2009	_	-	-	_	_	_	_	-	_	+	+	+	_	-

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Primary Study	Espallargues and Pons ¹⁶	Lo et al. ²⁴	Wang et al. ²⁶	Arrich et al. ¹⁰	Modawal et al. ²⁰	Strand et al. ¹⁷	Reichenbach et al. ²¹	Bellamy et al. ²²	Bannuru et al. ²³ (2009)	Rutjes et al. ²⁵	Colen et al. ¹⁸	Miller and Block ¹⁹	Chang et al. ²⁷	Bannuru et al. ¹¹ (2014)
Dixon, 1988	—	+	+	+	_	_	_	+	-	+	+	—	_	_
Dougados, 1993	-	+	+	+	-	_	_	+	_	+	+	—	_	-
Esteve de Miguel, 1995	_	-	_	-	_	-	_	-	_	+	_	_	_	_
Filardo, 2011	-	_	_	_	-	_	_	_	_	_	_	—	+	-
Filardo, 2012	-	_	_	_	-	_	_	_	_	_	_	—	+	-
Filardo, 2012a	-	_	_	_	-	_	_	_	_	_	_	—	+	-
Formiguera, 1995	—	-	+	_	_	_	_	+	_	+	_	_	—	—
Forster, 2003	-	_	_	_	-	_	_	+	_	_	+	—	_	-
Frizziero, 2002	-	_	_	_	-	_	_	+	+	_	+	—	_	-
Garcia, 2004	-	_	_	_	-	_	+	_	_	_	_	—	_	-
Genzyme, 2005	_	_	—	—	_	—	—	_	—	+	_	—	_	_
Ghirardini, 1990	—	-	_	_	_	_	_	_	_	+	_	_	—	—
Gobbi, 2012	_	_	_	_	_	_	_	_	_	_	_	_	+	_
Graf, 1993	_	_	_	_	-	_	_	+	_	_	+	_	_	_
Graf von der Schulenburg, 1997	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Grecomoro, 1987	_	_	+	+	+	_	_	+	_	+	+	+	_	-
Groppa, 2001	_	_	_	_	-	_	_	+	_	+	_	_	_	-
Groppa, 2004	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Guler, 1996	_	_	_	_	-	_	_	+	_	+	_	_	_	-
Halpern, 2013	_	_	_	_	_	_	_	_	_	_	_	_	+	_
Henderson, 1994	_	+	+	+	+	_	_	+	_	+	+	+	_	_
Heybeli, 2008	_	_	_	_	_	_	_	_	_	+	+	_	_	_
Hizmetli, 1999	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Hizmetli, 2002	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Huang, 2005	_	_	_	_	_	_	_	+	_	+	_	_	_	_
Huang, 2011	_	_	_	_	_	_	_	_	_	+	_	+	_	_
Huskisson, 1999	_	+	+	+	+	_	_	+	_	+	+	+	_	_
Isdale, 1993	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Ishjima, 2012	_	_	_	_	_	_	_	_	_	_	_	_	_	+
Jang, 2013	_	_	_	_	_	_	_	_	_	_	_	_	+	_
Jones, 1995	_	_	_	_	_	_	_	+	+	_	_	_	_	_
Jorgensen, 2010	_	_	_	_	_	_	_	_	_	+	+	+	_	_
Jubb, 2001	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Jubb, 2003	_	+	_	+	_	_	_	+	_	+	+	+	_	_
Juni, 2007	_	_	_	_	_	_	+	_	_	_	+	_	_	_
Kahan, 2001	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Kahan, 2003	_	_	_	_	_	_	_	+	_	+	+	_	_	_
Kalay, 1997	_	_	_	_	_	_	_	+	_	+	_	_	_	_
Karatay, 2004	_	_	_	_	_	_	+	+	_	_	_	_	_	_
Karatay, 2005	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Karatosun, 2005	_	_	_	_	_	_	+	+	_	_	+	_	_	_
Karlsson, 1999	_	_	_	_	_	_		+	_	_	_	_	_	_
Karlsson, 2002	_	+	_	+	_	_	+	+	_	+	+	+	_	_
Karlsson, 2003	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Karras, 2001	_	_	_	_	_	—	—	+	—	—	_	_	_	_

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Primary Study	Espallargues and Pons ¹⁶	Lo et al. ²⁴	Wang et al. ²⁶	Arrich et al. ¹⁰	Modawal et al. ²⁰	Strand et al. ¹⁷	Reichenbach et al. ²¹	Bellamy et al. ²²	Bannuru et al. ²³ (2009)	Rutjes et al. ²⁵	Colen et al. ¹⁸	Miller and Block ¹⁹	Chang et al. ²⁷	Bannuru et al. ¹¹ (2014)
Kawabata, 1993	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Kawasaki, 2009	-	-	_	_	-	_	—	-	-	_	+	_	_	-
Kirchner, 2005	-	-	_	_	-	_	—	+	-	_	_	_	_	-
Kirchner, 2006	-	-	_	_	-	_	+	+	-	_	+	_	_	-
Kon, 2010	-	-	_	_	-	_	—	-	-	_	_	_	+	-
Kon, 2011	-	-	_	_	-	_	—	-	-	_	_	_	+	-
Kosuwon, 2010	—	_	_	—	—	—	_	-	_	+	—	_	—	—
Kotevoglu, 2002	-	-	_	_	-	_	—	+	-	_	_	_	_	-
Kotevoglu, 2005	-	-	_	_	-	_	—	+	-	_	_	_	_	-
Kotevoglu, 2006	-	-	_	_	-	_	+	-	-	+	+	+	_	-
Kul-Panza, 2010	—	-	_	_	_	_	_	-	_	+	+	+	_	—
Lanzer, 2002	-	-	_	_	-	_	—	+	-	_	_	_	_	-
Leardini, 1987	-	-	_	_	-	_	—	+	+	_	+	_	_	-
Leardini, 1991	—	_	_	_	—	—	_	+	+	—	+	_	—	—
Lee, 2006	—	-	_	_	_	_	_	-	_	_	+	—	_	—
Lee, 2011	—	_	_	_	—	—	_	-	_	—	+	_	—	—
Leopold, 2003	—	-	_	_	_	_	_	+	_	_	+	—	_	—
Li, 2011	—	_	_	_	—	—	_	-	_	—	—	_	+	—
Lin, 2004	—	-	_	_	_	_	_	+	_	_	_	—	_	—
Listrat, 1997	—	_	_	_	—	—	_	+	_	+	—	_	—	—
Lohmander, 1996	_	+	+	+	+	+	_	+	_	+	+	+	_	_
Lundsgaard, 2008	—	_	_	—	—	—	_	-	_	+	+	+	—	—
Lussier, 1996	+	_	_	_	-	_	_	_	-	_	_	_	_	_
Marshall, 1999	+	_	_	—	—	—	_	-	_	—	—	_	—	—
McDonald, 2000	—	_	_	—	—	—	_	+	_	—	—	_	—	—
Miller, 1999	+	-	_	_	-	_	—	-	-	_	_	_	_	-
Miltner, 2002	—	-	_	_	_	_	_	+	_	+	_	—	_	—
Moreland, 1993	+	_	_	—	—	—	_	+	_	+	—	_	—	—
Nahler, 1996	-	-	_	_	-	_	—	+	-	_	_	_	_	-
Nahler, 1998	—	-	_	_	_	_	_	+	_	_	_	—	_	—
Napolitano, 2012	-	-	_	_	-	_	—	-	-	_	_	_	+	-
Navarro-Sarabia, 2011	-	-	_	_	-	_	—	-	-	+	_	_	_	-
Neustadt, 2004	_	-	-	-	_	-	_	+	_	+	-	_	-	-
Neustadt, 2005	-	-	_	_	-	_	—	+	-	_	+	_	_	-
O'Hanlon, 1995	+	-	-	-	_	-	_	-	_	-	-	_	-	-
Onel, 2008	-	-	_	_	-	_	—	-	-	_	+	_	_	-
Ozturk, 2005	—	-	_	_	_	_	_	+	_	_	_	—	_	—
Ozturk, 2006	-	-	_	_	-	_	—	-	-	_	+	_	_	-
Patel, 2013	—	-	_	_	_	_	_	-	_	_	_	—	+	—
Patrella, 2002	—	-	_	_	_	_	_	-	_	_	+	—	_	—
Pavelka, 2010	—	_	—	—	—	—	_	_	_	+	—	—	—	—
Payne, 2000	_	_	_	_	_	_	_	-	-	+	_	—	_	—
Pedersen, 1993	—	_	—	—	—	—	_	_	_	+	—	—	—	—
Petrella, 2002	—	+	—	+	+	—	_	+	_	+	—	—	—	+
Petrella, 2006	_	-	-	_	-	_	_	-	_	+	+	_	_	_

ARTICLE IN PRESS VISCOSUPPLEMENTATION TREATMENT FOR KNEE OA

Primary Study	Espallargues and Pons ¹⁶	Lo et al. ²⁴	Wang et al. ²⁶	Arrich et al. ¹⁰	Modawal et al. ²⁰	Strand et al. ¹⁷	Reichenbach et al. ²¹	Bellamy et al. ²²	Bannuru et al. ²³ (2009)	Rutjes et al. ²⁵	Colen et al. ¹⁸	Miller and Block ¹⁹	Chang et al. ²⁷	Bannuru et al. ¹¹ (2014)
Petrella, 2008	_	_	_	_	_	_	_	_	_	+	+	+	_	_
Petrella, 2009	—	_	—	—	—	—	_	_	_	+	—	_	—	—
Pham, 2003	_	+	_	_	_	_	_	+	_	_	_	_	_	—
Pham, 2004	_	_	_	_	_	_	_	+	_	+	+	—	_	_
Pietrogrande, 1991	—	_	—	_	—	—	_	+	+	—	+	_	—	—
Puhl, 1993	_	+	+	+	+	+	_	+	_	+	+	+	_	_
Puttick, 1995	+	_	_	_	_	_	_	_	_	_	_	_	_	_
Raman, 2006	_	_	_	_	_	_	+	_	_	_	_	—	_	_
Raman, 2008	_	_	_	_	_	_	_	_	_	_	+	_	_	_
Raynauld, 1999	+	_	_	_	_	_	_	_	_	_	_	_	_	_
Raynauld, 2002	_	_	_	_	_	_	_	+	_	+	+	_	_	_
Raynauld, 2005	_	_	_	_	_	_	_	_	_	_	+	_	_	_
Redd, 2003	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Rejaili, 2005	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Renklitepe, 2000	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Rolf, 2005	_	_	_	_	_	_	+	_	_	_	_	+	_	_
Roman, 2000	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Russell, 1992	_	+	_	+	_	_	_	_	_	+	_	_	_	_
Rvdell, 1972	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Sala, 1995	_	+	_	+	_	_	_	_	_	_	_	+	_	_
Sampson, 2011	_	_	_	_	_	_	_	_	_	_	_	_	+	_
Sanchez, 2012	_	_	_	_	_	_	_	_	_	_	_	_	+	_
Sanofi-Aventis, 2010	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Saravanan, 2002	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Scale, 1994	+	+	+	+	+	_	_	+	_	+	+	+	_	_
Schneider, 1997	_	_	_	_	_	_	_	+	_	+	_	_	_	_
Seikagaku, 2001	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Seikagaku, 2001a	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Sezgin, 2005	_	_	_	_	_	_	_	+	_	+	_	_	_	_
Shichikawa, 1983	_	_	_	_	_	_	_	+	_	+	_	_	_	_
Shichikawa, 1983a	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Shimizu, 2010	_	_	_	_	_	_	_	_	_	_	+	_	_	_
Skwara, 2009	_	_	_	_	_	_	_	_	_	_	+	_	_	_
Skwara, 2009a	_	_	_	_	_	_	_	_	_	_	+	_	_	_
Spakova, 2012	_	_	_	_	_	_	_	_	_	_	_	_	+	_
Sripada, 1999	+	_	_	_	_	_	_	_	_	_	_	_	_	_
Stittik 2007	_	_	_	_	_	_	_	_	_	_	+	_	_	_
Strand 2012	_	_	_	_	_	_	_	_	_	_	_	+	_	_
Tamir 2001	_	+	+	+	_	_	_	+	_	+	+	_	_	_
Tascioglu 2003	_	_	_	_	_	_	_	+	+	_	+	_	_	_
Tekeoglu 1998	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Tetik 2003	_	_	_	_	_	_	_	_	_	+	_	_	_	_
Thompson 2002	_	_	_	_	_	_	_	+	_	_	_	_	_	_
Torrance 1999	Т.	_	_	_	_	_	_	_	_	_	_	_	_	_
Teai 2003	т _	_	_	_	_	_			_		_		_	_
1501, 2003	—	—	_	_	_	_	—	+	—	+	_	—	_	—

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Primary Study	Espallargues and Pons ¹⁶	Lo et al. ²⁴	Wang et al. ²⁶	Arrich et al. ¹⁰	Modawal et al. ²⁰	Strand et al. ¹⁷	Reichenbach et al. ²¹	Bellamy et al. ²²	Bannuru et al. ²³ (2009)	Rutjes et al. ²⁵	Colen et al. ¹⁸	Miller and Block ¹⁹	Chang et al. ²⁷	Bannuru et al. ¹¹ (2014)
Tsukamoto, 1995	_	_	_	_	_	_	_	+	_	_	_	_	_	-
Ulucay, 2007	_	_	_	_	—	—	_	—	_	—	+	_	_	_
Vanelli, 2010	_	_	_	_	—	—	_	—	_	—	+	_	_	_
Wang-Saegusa, 2011	_	_	_	_	_	_	_	_	-	_	_	_	+	_
Weiss, 1981	_	_	_	_	—	—	_	—	_	+	+	_	_	_
Weiss, 1981a	_	_	_	_	_	_	_	_	-	+	_	_	_	_
Weiss, 1999	+	_	_	_	_	_	_	_	_	_	_	_	_	-
Westrich, 2009	_	_	_	_	_	_	_	_	-	_	+	_	_	_
Wobig, 1998	+	+	+	+	+	_	_	+	_	+	+	+	_	-
Wobig, 1999	_	_	_	_	_	_	+	+	-	_	+	_	_	_
Wu, 1997	_	_	+	+	_	_	_	+	-	+	_	_	_	_
Wu, 2004	_	_	_	_	_	_	_	+	-	+	_	+	_	_
Yamamoto, 1994	_	_	_	_	_	_	_	+	-	_	_	_	_	_
Yentur, 2003	_	_	_	_	_	_	_	+	_	_	_	_	_	-
Zhou, 2000	_	_	_	_	—	_	+	_	-	_	_	—	_	—

NOTE. The designation "a" after the date of a primary study indicates a separate study from the same author and the same calendar year.

	
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Appendix Table 2	. Heterogeneity c	or Subgroup	Analyses	of Primary Studies
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									Bannuru					Bannuru
	Espallargues and Pons ¹⁶	Lo et al. ²⁴	Wang et al. ²⁶	Arrich et al. ¹⁰	Modawal et al. ²⁰	Strand et al. ¹⁷	Reichenbach et al. ²¹	Bellamy et al. ²²	et al. ²³ (2009)	Rutjes et al. ²⁵	Colen et al. ¹⁸	Miller and Block ¹⁹	Chang et al. ²⁷	et al. ¹¹ (2014)
Statistical heterogeneity analysis	_	+	+	+	+	_	+	+	+	+	+	+	+	+
Subgroup or statistical analysis														
Pooled effect size, IA-HA v IA-placebo: after		+												
removal of highest-MW formulation HA														
Pain outcome, IA-HA v IA-placebo: in large trials										+				
Pain outcome, IA-HA v IA-placebo: in										+				
unpublished trials														
Pain outcome, IA-HA <i>v</i> IA-placebo: in blinding of outcome										+				
Pain outcome, IA-HA v IA-placebo: in large trials										+				
with blinding of outcome														
Physical function, IA-HA v IA-placebo: in large trials with blinding of outcome										+				
Pooled effect size, IA-HA <i>v</i> PRP: excluding all but													+	
RCTs														
Pooled effect size, IA-HA <i>v</i> NSAIDs: restricted to double-blinded studies														+
Pooled effect size IA-HA v IA-corticosteroids: in									+					
trials using intention to treat														
Pooled effect size IA-HA v IA-corticosteroids: in									+					
trials reporting single- or double-blind methodology														
Pain by VAS scale, IA-HA ν IA-placebo: good-					+									
Quality studies														
Kanuoni-enecis regression model, IA-HA: Dased					+									
On pain, with activity														
Random-enecis regression model, IA-HA: based					+									
On HA type—nyaluronan v nylan G-F 20														
Random-enecis regression model, IA-HA: based					+									
on study quality—poor v good														
rain at rest, IA-HA v IA-placedo: high-quality trials				+										
Pain during or after exercise, IA-HA ν IA- placebo: high-quality trials				+										
Function, IA-HA v IA-placebo: high-quality trials				+										
Pain with activities, IA-HA: cross-linked HA v			0	I										
non-cross-linked HA triais			0											
Function, IA-HA: cross-linked HA v non-cross- linked HA trials			0											
Major adverse events, IA-HA: non-cross-linked HA trials			0											
Major adverse events, IA-HA: cross-linked HA			0											
Pain with activities, IA-HA: non–cross-linked trials			+											

									Bannuru	l				Bannuru
	Espallargues	Lo	Wang	Arrich	Modawal	Strand	Reichenbach	Bellamy	et al. ²³	Rutjes	Colen	Miller and	Chang	et al. ¹¹
	and Pons ¹⁶	et al. ²⁴	et al. ²⁶	et al. ¹⁰	et al. ²⁰	et al. ¹⁷	et al. ²¹	et al. ²²	(2009)	_et al. ²⁵	et al. ¹⁸	Block	et al. ²⁷	(2014)
Pain with activities, IA-HA: non-cross-linked, single-blind trials			+											
Pain with activities, IA-HA: non-cross-linked, double-blind trials			+											
Pain with activities, IA-HA: non-cross-linked,			+											
single-center trials														
Pain with activities, IA-HA: non–cross-linked,			+											
multicenter trials														
Pain with activities, IA-HA: non-cross-linked, no			+											
intention-to-treat trials														
Pain with activities, IA-HA: non-cross-linked,			+											
only intention-to-treat trials														
Pain with activities, IA-HA: non-cross-linked			+											
trials with no escape analgesics														
Pain with activities, IA-HA: non-cross-linked			+											
trials with acetaminophen as escape analgesic														
Pain with activities, IA-HA: non-cross-linked, no			+											
restriction on escape analgesics														
Pain with activities, IA-HA: non-cross-linked,			+											
mean age of patients ≤ 65 yr														
Pain with activities, IA-HA: non-cross-linked,			+											
mean age of patients >65 yr														
Pain with activities, IA-HA: non–cross-linked			+											
trials without most advanced OA stage														
Pain with activities, IA-HA: non–cross-linked			+											
trials with most advanced OA stage														
Pain with activities, IA-HA: non–cross-linked, no			+											
restrictions on most advanced OA stage														
Pain with activities, IA-HA: non-cross-linked			+											
trials with entirities. In the new more links d														
rials with activities, IA-HA: non-cross-linked			+											
Pain with activities IA HA: non- gross linked no														
restriction on effusion criteria in trials			Ŧ											
Pain with activities IA HA: non-cross linked			1											
trial duration ≤ 12 wk			Ŧ											
Pain with activities $IA-HA$: non-cross-linked			+											
trial duration >12 wk			I											
Pain with activities IA-HA: non-cross-linked			+											
trials with sample size <100			1											
Pain with activities. IA-HA: non-cross-linked			+											
trials with sample size >100														
Pain with activities, IA-HA: non-cross-linked			+											
non-industry-funded trials														
Pain with activities, IA-HA: non-cross-linked,			+											
industry-funded trials														

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									Bannuru					Bannuru
	Espallargues	Lo	Wang	Arrich	Modawal	Strand	Reichenbach	Bellamy	et al. ²³	Rutjes	Colen	Miller and	Chang	et al. ¹¹
	and Pons ¹⁶	et al. ²⁴	et al. ²⁶	et al. ¹⁰	et al. ²⁰	et al. ¹⁷	et al. ²¹	et al. ²²	(2009)	et al.25	et al. ¹⁸	Block ¹⁹	et al. ²⁷	(2014)
Pain without activities, IA-HA: non-cross-linked			+											
trials														
Pain without activities, IA-HA: non–cross-			+											
linked, single-blind trials														
Pain without activities, IA-HA: non–cross-			+											
linked, double-blind trials														
Pain without activities, IA-HA: non–cross-			+											
linked, single-center trials														
Pain without activities, IA-HA: non–cross-			+											
linked, multicenter trials														
Pain without activities, IA-HA: non-cross-linked			+											
trials with no intention-to-treat analysis														
Pain without activities, IA-HA: non-cross-linked			+											
trials with intention-to-treat analysis														
Pain without activities, IA-HA: non-cross-linked			+											
trials with no escape analgesics														
Pain without activities, IA-HA: non-cross-linked			+											
trials with acetaminophen as escape analgesic														
Pain without activities, IA-HA: non-cross-linked			+											
trials with mean age of patients \leq 65 yr														
Pain without activities, IA-HA: non-cross-linked			+											
trials with mean age of patients >65 yr														
Pain without activities, IA-HA: non-cross-linked			+											
trials without most advanced OA stage														
Pain without activities, IA-HA: non-cross-linked			+											
trials with most advanced OA stage														
Pain without activities, IA-HA: non-cross-			+											
linked, no restrictions on most advanced OA														
stage														
Pain without activities, IA-HA: non-cross-linked			+											
trials with effusion as inclusion criteria														
Pain without activities, IA-HA: non-cross-linked			+											
trials with effusion as exclusion criteria														
Pain without activities, IA-HA: non-cross-			+											
linked, no restriction on effusion criteria in														
trials														
Pain without activities, IA-HA: non-cross-			+											
linked, trial duration ≤ 12 wk														
Pain without activities, IA-HA: non-cross-			+											
linked, trial duration >12 wk														
Pain without activities, IA-HA: non-cross-			+											
linked, non-industry-funded trials														
Pain without activities, IA-HA: non-cross-			+											
linked, industry-funded trials														
Functioning, IA-HA: non-cross-linked trials			+											

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									Bannuru					Bannuru
	Espallargues	Lo	Wang	Arrich	Modawal	Strand	Reichenbach	Bellamy	et al. ²³	Rutjes	Colen	Miller and	Chang	et al. ¹¹
	and Pons ¹⁶	et al. 24	et al. 26	et al. ¹⁰	et al. ²⁰	et al. ¹⁷	et al. ²¹	et al. ²²	(2009)	et al. ²⁵	et al. ¹⁸	Block	et al. ²⁷	(2014)
Functioning, IA-HA: non-cross-linked trials			+											
with no intention-to-treat analysis														
Functioning, IA-HA: non-cross-linked trials			+											
with intention-to-treat analysis														
Functioning, IA-HA: non-cross-linked trials			+											
with no escape analgesics														
Functioning, IA-HA: non-cross-linked trials			+											
with acetaminophen as escape analgesic														
Functioning, IA-HA: non-cross-linked trials			+											
with no restriction on escape analgesics														
Functioning, IA-HA: non-cross-linked trials			+											
without most advanced OA stage														
Functioning, IA-HA: non–cross-linked, no			+											
restrictions on most advanced OA stage														
Functioning, IA-HA: non-cross-linked trials			+											
with effusion as inclusion criteria														
Functioning, IA-HA: non–cross-linked trials			+											
with effusion as exclusion criteria														
Functioning, IA-HA: non–cross-linked, no			+											
restriction on effusion criteria in trials														
Functioning, IA-HA: non–cross-linked, trial			+											
duration ≤ 12 wk														
Functioning, IA-HA: non–cross-linked, trial			+											
duration >12 wk														
Functioning, IA-HA: non–cross-linked, non			+											
—industry-funded trials														
Functioning, IA-HA: non–cross-linked, industry-			+											
Pain outcome IA HA v IA placebo: trials with										I				
adequate concealment of allocation										Ŧ				
Pain outcome IA-HA v IA-placebo: trials with										_L				
inadequate concealment of allocation														
Pain outcome IA-HA v IA-placebo: trials with										+				
adequate v inadequate concealment of										I				
allocation														
Pain outcome IA-HA ν IA-placebo: trials with										+				
sham intervention										I				
Pain outcome, IA-HA ν IA-placebo: trials with										+				
non-sham intervention														
Pain outcome, IA-HA v IA-placebo: trials with										+				
sham v non-sham intervention														
Pain outcome, IA-HA v IA-placebo: trials with										+				
adequate blinding of patients														
Pain outcome, IA-HA v IA-placebo: trials with										+				
inadequate blinding of patients														

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	Espallargues	Lo	Wang	Arrich	Modawal	Strand	Reichenbach	Bellamy	et al. ²³	Rutjes	Colen	Miller and	Chang	et al. ¹¹
	and Pons ¹⁶	et al. ²⁴	et al. ²⁶	et al. ¹⁰	et al. ²⁰	et al. ¹⁷	et al. ²¹	et al. ²²	(2009)	et al.23	et al. ¹⁸	Block	et al. ²⁷	(2014)
Pain outcome, IA-HA v IA-placebo: trials with										+				
Pain outcome IA-HA v IA-placebo: trials with										-				
adequate blinding of outcome assessment										T				
Pain outcome IA-HA v IA-placebo trials with										+				
inadequate blinding of outcome assessment														
Pain outcome, IA-HA v IA-placebo: trials with										+				
adequate <i>v</i> inadequate blinding of outcome assessment														
Pain outcome, IA-HA v IA-placebo: trials with										+				
intention-to-treat analysis														
Pain outcome, IA-HA v IA-placebo: trials with no intention-to-treat analysis										+				
Pain outcome, IA-HA v IA-placebo: trials with										+				
intention-to-treat v no intention-to-treat analysis														
Pain outcome, IA-HA v IA-placebo: trials with >100 patients										+				
Pain outcome. IA-HA v IA-placebo: trials with										+				
<100 patients														
Pain outcome, IA-HA v IA-placebo: trials with										+				
$\geq 100 v < 100$ patients														
Pain outcome, IA-HA v IA-placebo: full journal										+				
publications														
Pain outcome, IA-HA v IA-placebo: other										+				
publications														
Pain outcome, IA-HA v IA-placebo: unpublished										+				
studies														
Pain outcome, IA-HA v IA-placebo: trials based										+				
on publication status														
rain outcome, IA-HA V IA-placebo: trials with										+				
Bain outcome IA HA : IA placebo: trials with														
industry-dependent funding										Ŧ				
Pain outcome IA-HA v IA-placebo: trials with										+				
industry-independent <i>v</i> industry-dependent										I				
funding														
Pain outcome. IA-HA v IA-placebo: trials with >1										+				
cycle of HA										'				
Pain outcome, IA-HA ν IA-placebo: trials with 1										+				
cycle of HA										·				
Pain outcome, IA-HA <i>v</i> IA-placebo: trials with >1										+				
v l cycle of HA														
Pain outcome, IA-HA v IA-placebo: trials with										+				
1-2 injections of HA														

									Bannuru					Bannuru
	Espallargues	Lo	Wang	Arrich	Modawal	Strand	Reichenbach	Bellamy	et al. ²³	Rutjes	Colen	Miller and	Chang	et al. ¹¹
	and Pons ¹⁰	et al.24	et al. ²⁰	et al. ¹⁰	et al. ²⁰	et al."	et al. ²¹	et al. ²²	(2009)	et al. ²⁹	et al. ¹⁸	Block	et al. ²⁷	(2014)
Pain outcome, IA-HA v IA-placebo: trials with 3										+				
injections of HA														
Pain outcome, IA-HA <i>v</i> IA-placebo: trials with >3										+				
injections of HA														
Pain outcome, IA-HA v IA-placebo: trials based										+				
on No. of injections of HA														
Pain outcome, IA-HA v IA-placebo: trials with										+				
cross-linked HA														
Pain outcome, IA-HA v IA-placebo: trials with										+				
non-cross-linked HA														
Pain outcome, IA-HA v IA-placebo: trials based										+				
on structure of HA														
Pain outcome, IA-HA V IA-placedo: triais with										+				
Very nign—MW HA														
pain outcome, IA-HA V IA-placedo: thais with										+				
Dain outcome IA HA u IA placebou trials with														
low MW HA										Ŧ				
Pain outcome IA-HA v IA-placebo: trials based										1				
on MW of HA										T				
Pain outcome IA-HA ν IA-placebo: trials with										+				
follow-up >6 mo										1				
Pain outcome. IA-HA v IA-placebo: trials with										+				
follow-up of 3-6 mo														
Pain outcome, IA-HA v IA-placebo: trials with										+				
follow-up <3 mo														
Pain outcome, IA-HA v IA-placebo: trials based										+				
on follow-up duration														
Pain-related outcomes, hylan v IA-HA: trials with							+							
adequate concealment of allocation														
Pain-related outcomes, hylan v IA-HA: trials with							+							
inadequate or unclear concealment of														
allocation														
Pain-related outcomes, hylan v IA-HA: trials with							+							
inadequate or unclear v adequate concealment														
of allocation														
Pain-related outcomes, hylan v IA-HA: trials with							+							
blinding of patients														
Pain-related outcomes, hylan v IA-HA: trials							+							
without blinding of patients														
Pain-related outcomes, hylan v IA-HA: trials with							+							
blinding ν without blinding of patients														
Pain-related outcomes, hylan v IA-HA: trials with							+							
intention-to-treat analysis														

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									Bannuru					Bannuru	
	Espallargues	Lo	Wang	Arrich	Modawal	Strand	Reichenbach	Bellamy	et al. ²³	Rutjes	Colen	Miller and	Chang	et al.11	
	and Pons ¹⁶	et al. ²⁴	et al. ²⁶	et al. ¹⁰	et al. ²⁰	et al. ¹⁷	et al. ²¹	et al. ²²	(2009)	et al. ²⁵	et al. ¹⁸	Block ¹⁹	et al. ²⁷	(2014)	
Pain-related outcomes, hylan v IA-HA: trials							+								
without intention-to-treat analysis															
Pain-related outcomes, hylan v IA-HA: trials with							+								
intention-to-treat v no intention-to-treat															
Pain-related outcomes hylan v [A-HA: trials with							+								
No. of patients randomized >200							I								
Pain-related outcomes, hylan v IA-HA: trials with							+								
No. of patients randomized ≤ 200															
Pain-related outcomes, hylan v IA-HA: trials with							+								
No. of patients randomized >200 $\nu \leq 200$															
Pain-related outcomes, hylan v IA-HA: trials with							+								
follow-up >3 mo															
Pain-related outcomes, hylan v IA-HA: trials with							+								
follow-up $\leq 3 \mod 3$															
Pain-related outcomes, hylan <i>v</i> IA-HA: trials with							+								
follow-up >3 mo $\nu \leq 3$ mo															
Pain-related outcomes, hylan v IA-HA: large							+								
trials															
Pain-related outcomes, nylan v IA-HA: small trials							+								
Pain-related outcomes, hylan v IA-HA: trial							+								
comparison of interaction based on size															
Flare-ups, IA-HA ν IA-placebo: large trials with										+					
blinded outcome															
Serious adverse events, IA-HA v IA-placebo:										+					
large trials with blinded outcome															
Dropouts due to adverse events, IA-HA v IA-										+					
placebo: large trials with blinded outcome															
Overall adverse events, IA-HA v IA-placebo: large										+					
trials with blinded outcome															
Effusions, IA-HA v IA-placebo: large trials with										+					
blinded outcome															
Local adverse events, IA-HA v IA-placebo: large										+					
trials with blinded outcome															
Overall study withdrawals, IA-HA v IA-placebo:										+					
large trials with blinded outcome															

NOTE. A plus sign indicates formal sensitivity or subgroup analysis was performed, a minus sign indicates formal sensitivity or subgroup analysis was not performed, and a zero indicates descriptive data were provided or discussed but no analysis was performed.

HA, hyaluronic acid; IA, intra-articular; MW, molecular weight; NSAIDs, nonsteroidal anti-inflammatory drugs; OA, osteoarthritis; PRP, platelet-rich plasma; RCTs, randomized controlled trials; VAS, visual analog scale.

	Espallargues and Pons ¹⁶	Lo et al. ²⁴	Wang et al. ²⁶	Arrich et al. ¹⁰	Modawal et al. ²⁰	Strand et al. ¹⁷	Reichenbach et al. ²¹	Bellamy et al. ²²	Bannuru et al. ²³ (2009)	Rutjes et al. ²⁵	Colen et al. ¹⁸	Miller and Block ¹⁹	Chang et al. ²⁷	Bannuru et al. ¹¹ (2014)
ІА-НА									/					
Knee function												+		
Improvement in symptoms	0													
No requirement for TKA	0													
Knee pain outcomes												+		
Adverse events	0		+							+				
Mortality			0											
IA-HA (including specific HA products) v IA-	-placebo													
Overall pooled effect size	1	+												
Lequesne index score						+		+						
Knee function	+			+				+				+		
Physical function										+				
Painful symptoms of knee OA (WB pain)	+													
Pain with activities			+					+						
Pain during or immediately after exercise				+				+						
Patient global assessment								+						
Knee circumference								Ó						
Pain at rest				+				+						
Percentage of painful days				'				+						
Knee pain outcomes					+	0		+		+	+	+		
WOMAC scores						Ū		+			'	I.		
Overall adverse events				+				+		+		+		
Flare-ups				'				'		+		I.		
Systemic reactions								+						
Injection-site reaction						0		'						
Injection-site pain						0								
Arthralgia						0								
Arthropathy/arthrosis/arthritis						0								
Back pain						0								
Headache						0								
Knee effusion						0				1				
Discontinued due to adverse event						0				т 				
Overall study withdrawal						0		+		Т				
No. of clinical failures								+						
No. of survivors								+						
Knee ROM								- -						
Joint space width								т 						
IA-HA 1/ oral NSAIDs								Г						
Overall pooled effect size	Т													1
Knee pain outcomes	T							0						一 一
WOMAC scores								0						T
Knee stiffness								U						J
Serious adverse events														+
Local adverse events								0						0
								U						
													(continued)

Appendix Table 3. Outcomes That Were Assessed for and Reported by Included Studies

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	Espallargues and Pons ¹⁶	Lo et al. ²⁴	Wang et al. ²⁶	Arrich et al. ¹⁰	Modawal et al. ²⁰	Strand et al. ¹⁷	Reichenbach et al. ²¹	Bellamy et al. ²²	Bannuru et al. ²³ (2009)	Rutjes et al. ²⁵	Colen et al. ¹⁸	Miller and Block ¹⁹	Chang et al. ²⁷	Bannuru et al. ¹¹ (2014)	.el4
Study withdrawal							_	0							
Injection-site pain								0							
Injection-site swelling								0							
IA-HA v IA-PRP															
Overall pooled effect size													+		
Risk of adverse reactions													+		
IA-HA (various specific products) v IA-cortic	osteroids (vari	ous prepa	arations)												
Overall pooled effect size									+						
Knee pain outcomes								+							
Total Larson rating score								0							
Knee ROM								+							
Function								0							
Patient global assessment								+							
Study withdrawal								+							
Local adverse reactions								+							
Systemic adverse reactions								+							
Analgesic use								0							Κ.
Comparison of various specific IA-HA produ-	cts														А.
Knee pain outcomes							+	+			+				Ω
Patient global assessment								+							4M
WOMAC scores								0							IPE
Function								+							BEI
Overall pooled effect size	+						+				+				Ľ
Knee ROM								0							ET
Local adverse events							+	+							Ā
Flare-ups							+								L.
Joint effusion							+								
Painful injections								+							
Lequesne index score								0							
Clinical failures								0							
Study withdrawal								0							

NOTE. A plus sign indicates formal sensitivity or subgroup analysis was performed, and a zero indicates descriptive data were provided or discussed but no analysis was performed. HA, hyaluronic acid; IA, intra-articular; NSAIDs, nonsteroidal anti-inflammatory drugs; OA, osteoarthritis; PRP, platelet-rich plasma; ROM, range of motion; TKA, total knee arthroplasty; WB, weight bearing; WOMAC, Western Ontario and McMaster Universities Osteoarthritis Index.