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Hypertonic dextrose injections (prolotherapy) in the treatment of symptomatic knee osteoarthritis: A systematic review and metaanalysis

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Hypertonic dextrose injections (prolotherapy) is an emerging treatment for symptomatic knee osteoarthritis (OA) but its efficacy is uncertain. We conducted a systematic review with meta-analysis to synthesize clinical evidence on the effect of prolotherapy for knee OA. Fifteen electronic databases were searched from their inception to September 2015. The primary outcome of interest was score change on the Western Ontario and McMaster Universities Arthritis Index (WOMAC). Three randomized controlled trials (RCTs) of moderate risk of bias and one quasi–randomized trial were included, with data from a total of 258 patients. In the meta-analysis of two eligible studies, prolotherapy is superior to exercise alone by a standardized mean difference (SMD) of 0.81 (95% CI: 0.18 to 1.45, p=0.012), 0.78 (95% CI: 0.25 to 1.30, p=0.001) and 0.62 (95% CI: 0.04 to 1.20, p=0.035) on the WOMAC composite scale; and WOMAC function and pain subscale scores respectively. Moderate heterogeneity exists in all cases. Overall, prolotherapy conferred a positive and significant beneficial effect in the treatment of knee OA. Adequately powered, longer-term trials with uniform end points are needed to better elucidate the efficacy of prolotherapy.

Knee osteoarthritis (OA) is the most common form of chronic arthritis worldwide and a major cause of pain and disability¹⁻³. It carries considerable economic burden with its high prevalence, loss of work time and utilization of healthcare service resources^{4,5}. The effectiveness of current treatment for KOA is limited. Current guidelines suggest that the management of knee OA should take a multidisciplinary approach including non-pharmacological, pharmacological, surgical and complementary therapies^{6,7}. Despite efforts by the Osteoarthritis Research Society International (OARSI) Treatment Guidelines Committee, there are limited generally accepted core sets of treatments for patients with knee OA⁸. A safe and effective treatment option that complements the current therapy remains a top priority in clinical practice and research. In the United States, assessment of knee OA treatment has been identified as a "top 100" research priority by the National Academy of Medicine⁹, and the Agency for Healthcare Research and has called for new knee OA therapy¹⁰.

Hypertonic dextrose injection, "prolotherapy", is an injection-based treatment used for a variety of painful chronic musculoskeletal pain conditions, including knee OA¹¹. The core practice principle of prolotherapy is injection of relatively small volumes (0.5–6 ml) of an irritant solution, usually hypertonic dextrose, at painful ligament and tendon attachments, as well as in adjacent joint spaces¹¹. The basic science of the mechanism of action on connective tissue and cells of various type (e.g. fibroblast/chondrocytes etc.) is not well understood. However, animal studies have reported that peritendinous dextrose injection consistently resulted in fibroblast and vascular proliferation, dense collagen deposition and increase in ligament thickness, energy absorption and ultimate

¹The Chinese University of Hong Kong, The Jockey Club School of Public Health and Primary Care, Hong Kong. ²The University of Kansas, Department of Physical Medicine and Rehabilitation (1986-2015), Kansas City, USA. ³University of Wisconsin School of Medicine and Public Health, Department of Family Medicine and Community Medicine, Madison, Wisconsin, USA. Correspondence and requests for materials should be addressed to V.C.C. (email: vchunq@cuhk.edu.hk) load bearing ability^{12–15}. Animal model data has also suggested cartilage-specific anabolic growth as a result of intra-articular dextrose injection¹⁶. The hypothesized mechanisms for pain relief include: (i) stimulation of local healing among chronically injured extra- and intra-articular tissue; (ii) reduction of joint instability through the strengthening of stretched or torn ligaments, and (iii) stimulation of cellular proliferation¹⁷.

Prolotherapy is practiced throughout the world; the strongest interest appears to be among physicians and patients in primary care^{18,19}. Human studies have assessed the role of prolotherapy for various musculoskeletal conditions^{20–24}. Systematic reviews of the clinical effectiveness of prolotherapy in chronic low back pain and lateral epicondylitis had been conducted in the past which yielded mixed and positive results respectively, though the strength of evidence was limited by clinical heterogeneity amongst studies and the presence of co-interventions^{25,26}. In recent years, prolotherapy has been used to treat patients with knee OA refractory to other conservative care²⁷. Results from several published clinical trials have shown positive effects of prolotherapy in knee OA but the findings have not been synthesized^{28–31}. Therefore, we conducted a systematic review with the aim of more comprehensively assessing the efficacy of prolotherapy for knee OA in order to clarify its potential role as a non-surgical treatment modality.

Methods

We follow the PRISMA Reporting Guidelines for Systematic Review and Meta-analysis³²:

Search methods for identification of studies. Potential studies were identified by searching the electronic databases listed below, with search period starting from their inception till September 2015 (Appendix 1).

Cochrane Central Register of Controlled Trials (CENTRAL)

OVID MEDLINE

EMBASE

Global Health,

NHS Health Technology Assessment Database,

Digital Dissertation Consortium,

International Pharmaceutical Abstract

BIOSIS Preview.

AMED

Inspec

Ovid Nursing Database

CinicalTrial.gov (http://clinicaltrials.gov/),

Drugs@FDA (http://www.accessdata.fda.gov/scripts/cder/drugsatfda/index.cfm),

European Medicines Agency public assessment reports (EPAR,http://www.ema.europa.eu/ema),

Pharmaceuticals and medical devices agency of Japan (http://www.pmda.go.jp/english/service/approved. html).

Types of studies. This systematic review included randomized controlled trials (RCTs) and quasi-randomized controlled trials that compared prolotherapy injections to saline injections, water injections or exercise therapy. Co-interventions were allowed, as long as they were uniform across all groups. There were no limitations on publication dates. The protocol was registered at PROSPERO (CRD42015015901).

Types of participants. We selected studies that included participants aged 18 years or over with (i) a diagnosis of knee OA according to the criteria from the American College of Rheumatology³³, (ii) knee pain for at least 3 months and (iii) had reported the Western Ontario and McMaster Universities Arthritis Index (WOMAC)³⁴ or the Visual Analogue Scales (VAS) for pain as one of the outcomes³⁵. We excluded studies that included participants who had undergone total knee replacement.

Types of intervention. For inclusion, dextrose prolotherapy injections had to be administered to at least one group within the trial. For comparison groups we included studies that provided injections with 0.9% normal saline or water; or exercise therapy. Consistent with the clinical practice of prolotherapy, at least part of the injection protocol had to include an intra-articular injection, with or without additional injections to the peri-articular ligament or tendon attachments.

Outcome measures. In accordance with international consensus on the core set of outcome measures for phase III clinical trials in OA, eligible trials needed to include assessment of either self- reported pain or self-reported physical function³⁶. Following this recommendation, the primary outcome of interest is the Western Ontario and McMaster Universities Arthritis Index (WOMAC)³⁷. Secondary outcomes of pain include the Visual Analogue Scales (VAS) for pain³⁵, Knee Pain Scale (KPS)³⁸ and Wong-Baker Scale³⁹.

Eligibility Assessment and Data Extraction. Two reviewers (RS and VC) independently screened electronically retrieved titles and abstracts, evaluated potentially relevant full texts, and determined study eligibility. For eligible studies, data were extracted independently by the two authors (RS and DC) using a piloted data extraction form. For each eligible study, the following data were extracted: study design, clinical settings, participant characteristics, features of interventions, outcomes, duration of follow up and adverse events. An attempt was made to contact study authors regarding these methodological elements if not reported.

Risk of bias assessment. The risk of bias among included studies was assessed by the Cochrane's risk of bias tool⁴⁰ by two reviewers independently (XYW and RH). The following risk of bias domains were evaluated: sequence generation, allocation concealment, blinding of participants and research personnel, blinding of

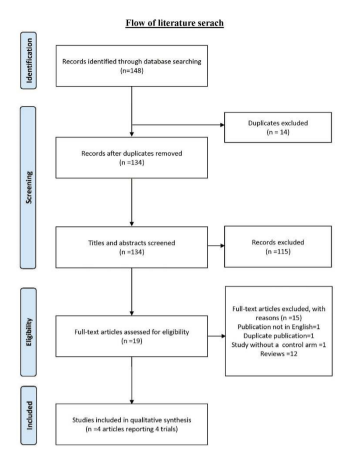


Figure 1. Flow of literature search.

outcome assessors, incomplete outcome data, and selective outcome reporting. Discrepancies in study selection, data extraction and risk of bias assessment results were resolved by group consensus. In the group consensus process, the two reviewers discussed reasons for discrepancy with a goal to achieve consensus after clarification. If a consensus could not be reached, a third reviewer Vincent Chung (VC) was included as an arbitrator. A decision was made after reevaluation of the included studies and further discussion between the three reviewers. This has been clarified in the revised manuscript.

Statistical Analysis. All meta-analyses were conducted using the STATA software⁴¹. A random effect model was used to pool study results, taking into account possible variations in effect sizes across trials⁴². Changes in continuous outcomes were pooled as standardized mean differences (SMD), as different scaling of outcome measurements across trials were expected. 95% confidence intervals (CI) were calculated for all estimates. I square (I^2) statistic was calculated to estimate heterogeneity across studies. An I^2 level of less than <25%, 25–50% and greater than 50% were regarded as indicators of low, moderate and high levels of heterogeneity respectively⁴³.

Minimal Clinically Important Difference (MCID) for WOMAC and VAS. The effect estimates were interpreted according to the values of the established Minimal Clinically Important Difference (MCID). It has been reported that a 12-point increment on WOMAC composite scale indicates 'good to very good' improvement when the scale is transformed onto a 0–100 point scale^{44,45}.

The MCID for VAS global pain assessment score and VAS pain on motion were based on a study in 2005; of which the MCID for improvement in the global VAS is 18.3 mm whereas that for VAS in motion is 19.9 mm⁴⁴.

Results

We identified 134 citations from all searches and excluded 14 duplicates. After screening the titles and abstracts, we retrieved 19 full texts for further assessment. Of these, 15 were excluded for the following reasons: duplicate publication as conference abstract (n = 1), publication not in English (n = 1), trial without a control arm (n = 1) and narrative reviews (n = 12). Four full texts which reported results from four clinical trials were eligible for inclusion^{28–31}.(Fig. 1)

Characteristics of included trials. Characteristics of included trials are summarized (Table 1). The four included clinical trials^{28–31} recruited a total of 258 patients with a diagnosis of knee OA based on American College of Rheumatology (ACR) guidelines including baseline radiological severity as graded by a Kellgren-Lawrence

Source & no. of arms	Interven-	Injection route and solution con- centration (%)	Total no of partici- pants	Number of case analyz- ed	Age +/— standard deviation	Sex Fe- male (%)	Weight +/- standard deviation	Body mass index (BMI)	No of injections received +/-standard deviation	Outcomes assessed	Trial duration (Time for end points assessment)
Raba- go ³⁰ (2 arms)	Dextrose OR Dextrose plus Mor- rhuate	Intra-articular dextrose 25%, Extra-articular dex- trose 15%, Sodium Morrhuate 5%	27	24	54.5+/-6.8	54.2	Not reported	25% of the subjects with BMI ≤25 37.5% of the subjects with BMI 26–30 37.5% of the subjects with BMI ≥31	4.5+/-0.9		52 weeks
										WOMAC composite -WOMAC-pain -WOMAC-stiff -WOMAC-function	(week 5,9,12,24,52)
	Saline OR Exercise	Intra-articular and extra-articular saline 0.9%	18	13	57.4+/-7.6	61.5	Not reported	30.8% of the subjects with BMI ≤25, 30.8% of the subjects with BMI 26–30, 38.5% of the subjects with BMI ≥31	4.3+/-0.8	Cartilage volume	(at week 52 only)
Raba- go ²² (3 arms)	Dextrose	Intra-articular dextrose 25%, Extra-articular dextrose 15%,	33	30	56.8+/-7.9	63	Not reported	33% of the subjects with BMI \leq 25, 20% of the subjects with BMI 25–30, 47% of the subjects with BMI, \geq 30	3.95+/-1	WOMAC composite WOMAC-pain WOMAC-stiff WOMAC-function KPS (each knee)	52 weeks (week 5,9,12,24,52)
	Saline	Intra-articular and extra-articular saline 0.9%	31	29	56.8+/-6.7	69	Not reported	28% of the subjects with BMI \leq 25, 38% of the subjects with BMI 25–30, 34% with BMI \geq 30	3.71+/-1.1		
	Exercise	N/A	33	31	56.4+/-7.0	68	Not reported	19% of the subjects with BMI ≤25, 39% of the subjects with BMI 25–30, 42% of the subjects with BMI ≥30	N/A		
Dumais et al. ²⁸ (2 arms with cross over)	Dextrose	Intra-articular dextrose 20%, Extra-articular dextrose 15%	21	18	57.3+/-12.6	38.9	90.1 (22)	32.2+/-7.2	3.7+/-0.7	WOMAC composite -WOMAC-pain -WOMAC-stiff -WOMAC-function Pain intensity	32 weeks (data col- lected at week 16 before cross- over; then reassessed at week 32)
	Exercise	N/A	24	18	56.2+/-10.9	55.6	92.4 (17.2)	34.3+/-5.7	N/A	Functional impair- ment Wong-Baker Descriptive Numer- ical VAS Combined pain score Timed UGT	
Reeves et al. ²¹ (2 arms)	Dextrose	Intra-articular dextrose10%	36	35	60.0+/-12.5	51.4	86.6 (19.2)	Not reported	3	VAS- rest VAS- walking VAS-stair use Swelling Buck- ling/2 mon Flexion range	24 weeks (assessed at week 24 only)
	Water	Intra-articular bac- teriostatic water	35	32	65.3+/-11.0	34.3	90.5(19.1)	Not reported	3		

Table 1. Characteristics of included studies. KEY: BMI = Body Mass Index. VAS = Visual Analogue Scale. UGT = Up and Go Test. WOMAC = Western Ontario and McMaster Universities Arthritis Index. KPS = Knee Pain Scale.

(K-L) score (Table 2)⁴⁶. The follow up period ranged from five weeks to one year post-enrollment. The injection protocols varied; two studies referred to an existing published injection protocol⁴⁷.

Risk of bias. Risk of bias amongst included studies was moderate overall (Table 3). None provided a detailed publicly accessible protocol of study methods including planned a *priori* statistical analysis methods (e.g. study protocols on trial registration websites such as ClinicalTrials.gov) that allowed assessment of selective outcome reporting, but three of them have provided a detailed analysis plan via personal communication^{29–31}. Sequence generation was considered to have a low risk of bias in only two studies, and two had a low risk of bias in allocation concealment. Two studies blinded both patients and investigators, but two did not. Two studies reported blinding of assessors but not the remaining two. Risk of bias incurred from incomplete outcome data varied, with drop-out rates ranged from 0% to 27.8%. Since three out of four trials did not provide trial protocol, this can be regarded as a major source of bias.

Effect of interventions. The effect of interventions with different comparison groups and follow up duration is summarized in Table 4. Dumais $et\ al.^{28}$ used WOMAC version constructed on a five point (0-4) Likert scale: composite (0-96), pain (0-20), stiffness (0-8) and function (0-68). Lower score indicated better knee related outcomes. Rabago $et\ al.^{30,31}$ used a WOMAC version constructed on the same five point (0-4) point scale) Likert scale. Each of the individual subscales was then converted to a 0-100 score scale, with higher score reflecting improvement.

Study	Interventions and controls	Kellgren Lawerence Grade 2 or above			
Reeves 2010	Dextrose	36			
	Normal saline	25			
		Kellgren-Lawerenc Grade 1 or 2(mild)	Kellgren-Lawerence grade 3 or 4 (moderate to severe)		
Dumais ²⁸	Dextrose	3	15		
	Exercise	2	16		
Rabago ²²	Dextrose	11	14		
	Normal Saline	12	9		
	Exercise	9	14		
Rabago ³⁰	Dextrose (Dextrose alone or dextrose + morrhuate) & Control (normal saline or exercise)	157	96		

Table 2. Distribution of OA knee severity grading among participants of included trials.

Dextrose Injection versus Exercise on WOMAC index. In this comparison, two RCTs (n = 97) were eligible for pooling using follow-up data from 12 to 16 weeks^{28,30}. Although the same WOMAC tool was used, different measurement scales were adopted in these two trials; therefore, SMDs were calculated in the random effect meta-analyses. Pooled results favored prolotherapy in terms of WOMAC composite and WOMAC function scores (Figs 2 and 3). The pooled SMD of WOMAC composite score is 0.81 (95% CI: 0.18 to 1.45, p = 0.012, $I^2 = 53.6\%$); whereas the SMD of WOMAC-function score is 0.78 (95% CI: 0.25 to 1.30, p = 0.001, $I^2 = 34.5\%$). For WOMAC pain (Fig. 4), pooled results indicated that the SMD is 0.62 (95% CI: 0.04 to 1.20, p = 0.035, $I^2 = 46.2\%$). While the p value indicates statistically significant result as defined by p < 0.05, the lower end of the 95% CI is very close to zero, suggesting the imprecision of possible treatment effect. Overall, sizes of the SMD indicated that dextrose injection provides substantially stronger improvement when compared to exercise, but the lower end of 95% CI has covered SMD values of small effect sizes (i.e. SMD ≤ 0.2)⁴⁸. Such uncertainty is also reflected in the low to moderate degree of heterogeneity among all three meta-analyses, given all with I² values ranging from 34.5% to 53.6%⁴⁹. Although it is not included in meta-analysis due to differences in data collection time, Rabago et al.²² also reported WOMAC outcomes at 26 and 52 week follow-up. Dextrose injections outperformed exercise on the composite and function-subscale score (26 and 52 weeks) and the pain-subscale score (26 weeks) respectively³⁰.

Dextrose versus Saline Injection on WOMAC index at 12 weeks and 52 weeks. In two trials, dextrose injections, either on their own³⁰ or mixed with sodium morrhuate³¹, were found to be superior to normal saline in improving WOMAC composite and subscale scores to levels above the minimal clinical important difference (MCID) at 12 and 52 weeks³⁵. (Table 4)

Pain intensity on VAS at 16 and 24 weeks after prolotherapy. At 16 weeks, Dumais *et al.* reported an improvement in the VAS global pain score of 29.7 points (SE: 4.57), above the MCID at 24 weeks, Reeves *et al.* also reported improvement in VAS pain during walking of 13.9 (SE: 3.10) and during stair climb of 13.7 (SE 3.2) 29 , which were slightly lower than the relevant MCID 44 . Pooling of the two trials was not possible as the VAS measured slightly different outcomes; the former represented a global assessment of pain whereas the latter measured pain level during movement.

Change in pain severity at 16 weeks and 24 weeks after prolotherapy. At 16 week, Dumais *et al.* reported statistically significant improvement in Wong Baker Scale scores after prolotherapy (adjusted mean change: 1.27, SE: 0.33, p = 0.003)²⁸. At 24 weeks, Rabago *et al.*²² also reported a statistically significant improvement in WOMAC pain scores (adjusted mean change $15.50 \pm \text{SE } 3.56$, p < 0.05) and on the KPS (adjusted mean change: 0.92, SE: 0, 25, p < 0.05). The pain reduction magnitude of active participants was significantly greater when compared with their control groups (p < 0.05)³⁰.

Side effects. All four trials monitored side effects and adverse effects. Only one trial reported self-limited bruises after both dextrose (n = 3) and saline injections (n = 5)³⁰. This was an expected side effect and deemed to be of minimal clinical relevance due to its transient nature

Discussion

Pooling data from two RCTs, we report that peri- and intra-articular hypertonic dextrose knee injections in three to five sessions have a statistically significant and clinically relevant effect in the improvement of WOMAC composite score, functional and pain subscale at 12 to 16 weeks compared to formal at-home exercise. Self-reported outcomes favoring prolotherapy are also observed in unpooled data when dextrose prolotherapy groups are compared with other control groups. The majority of the effect sizes are higher than the MCID, and the benefits were sustained up to 1 year. The risk of bias level in the included studies was moderate. Overall, prolotherapy injections appear to be safe, but no study was powered to detect rare adverse events.

While the overall direction of the effect is positive, some uncertainty of the effect size still exists due to the low to moderate heterogeneity and wide confidence intervals. In addition, this systematic review identified

Source	Sequence generation	Allocation concealment	Blinding of participants and researchers	Blinding of outcome assessment	Incomplete outcome data addressed	Selective outcome reporting
Reeves ²¹	Low. Random sequence was generated by random number table.	Unclear. Relevant information was not reported.	Low. Identical control solution was used	Low. Films were separated in different packets so reading 1 film would not influence reading of the next. X ray films were read by the chief investigator. A database coordinator loaded results onto the database.	High. Lost to follow up was 9/77 = 11.7% > 10%	Unclear. No proto- col was provided.
Dumais ²⁸	Unclear. Authors did not provide information on how random sequence was generated.	Low. Opaque sealed envelopes were used.	High. It was an open-labeled trial.	High. It was an open-labeled trial.	High. Lost to follow up was $3/21 = 14.3\%$ in group A. $6/24 = 25\%$ in group B. Both groups $>10\%$	Unclear. No proto- col was provided.
Rabago ²²	Low. Random sequence was generated by computer.	Low. Allocation concealment of injectant was achieved through the use of sealed opaque enve- lopes (personal communica- tion, Rabago 2015)	Low. Both active and control solutions looked similar.	Low. The injector, outcome assessor, principal investigator, and participants were blinded to injection group by preparation of syringes off site; blinding was formally assessed among injection participants and injector using questionnaire.	Low. No lost to follow-up cases.	Low. No protocol was provided,but analysis plan was clearly described.
Rabago ³⁰	High. Non-randomized study design.	High. Non-randomized study design.	High. Single-arm uncontrolled study/ at-home exercise "control" was used.	High. Primary outcome was self-reported QOL.	High. Lost to follow up was $3/27 = 11.1\%$ in intervention group. $5/18 = 27.8\%$ in control group. Both $>10\%$	Unclear. No proto- col was provided.

Table 3. Risk of bias assessment.

several limitations of existing trials. None of the included studies provided a publicly accessible protocol, therefore the potential for selective outcome reporting exists; however, the analyses are generally straightforward and well-reported. The sample size was small with only four trials eligible for review and only portions of two trials could be pooled in meta-analyses. Small sample size led to a wide confidence interval; thus the effect sizes reported may be imprecise. However, pooled results provide guidance for the sample size needed in future clinical studies 50 . Due to a lack of uniform longer-term follow up data across both studies, pooling of results could only be done with data collected between 12 to 16 week follow-up. Given that prolotherapy is hypothesized to work by healing and regeneration over several months 51,52 , the effects reported here may underestimate longer term benefits. A recent open label study reported progressive improvement in WOMAC scores for most study participants through 2.5 ± 0.6 years (range 1.6-3.5 years) of follow up after prolotherapy 27 . Finally, low to moderate quantified heterogeneity was reported in pooled analyses; this is likely attributable to multiple factors, including differences in patient characteristics, control treatment, study design, injection protocol methods, dextrose concentration, follow-up duration and outcomes assessment methods.

While the positive findings in the evaluated studies suggest efficacy, their limitations restrict a rigorous determination of efficacy and clinical utility. The current analysis suggests studies with the following characteristics can be considered in the future: 1) study duration of at least one year. Prolotherapy is hypothesized to be a regenerative injection therapy; serial monthly injections are typically performed to stimulate hypothesized tissue effects; 2) uniform self-reported outcomes and time points using the WOMAC outcome measures³⁷; 3) objectively determined physical function and health status outcomes such as hospitalization, nursing home placement and decline in health^{53,54}. Inclusion of objective assessment including magnetic resonance imaging³¹ and intra-articular and serum biomarkers⁵⁵. 4) a *priori* subgroup analysis to identify the phenotype of patients who respond most favorably to prolotherapy;5) because the performance of prolotherapy appears to be superior to saline and exercise, future studies may consider formal effectiveness designs using exercise or other non-injection matched control comparators. This avoids injection-related discomfort and risk among controls, saves study conduct time and cost, and offers a comparison relevant to clinical practice; 6) overall quality of life and cost effectiveness evaluation using the EuroQol-5D, a commonly used measure that allows comparison to other interventions⁵⁶; 7) determination of the MCID of WOMAC score specific to prolotherapy.

Investigation of prolotherapy is in an early stage and several treatment protocol issues warrant further study to determine optimal strategy, including dextrose concentration and volume, treatment frequency and duration, and specific utility of intra compared with extra-articular injections. Studies in this review used the approach of dextrose injections into the intra-articular joint space and the extra-articular soft tissue attachment. Recently, more superficial application of hypertonic dextrose has been reported to provide an analgesic effect in the treatment of a variety of peripheral neuropathic pain conditions^{57,58}. Sensorineural effects of dextrose have been hypothesized³¹. Given that the etiology of knee OA pain is multifactorial, future trials may consider adding superficial dextrose injection as part of the protocol. Concomitant basic science studies using *in vitro* and animal model methodologies are warranted to better elucidate the mechanism of action of dextrose in the context of musculoskeletal pathology. Finally, studies in this review used palpation guidance for injections; future investigators should also consider the use of ultrasound guidance to improve injection accuracy and precision, limiting procedural variability⁵⁹.

Strengths and weaknesses of this systematic review. The main limitation of this systematic review is the limited number of studies and their relatively small sample size. We considered whether or not to perform a meta-analysis in addition to systematic narrative review, given that the small sample sizes of included trials might generate an unstable pooled effect size⁶⁰. A meta-analyses is included and resulted in increased statistical

VOMAC C	Compos	ite at 12–16 v	veek				
n	n	Scale Range ^	Pre-treat- ment mean score	Standard deviation	Post-treatment adjusted mean change	Standard error	Duration of follow up
	18	0-96	44.4	13.7	-21.8 €	2.95	16 week
	18	0-96	36.2	16.8	-6.1 €	3.28	
	30	0-100	63.1	15.0	13.31	3.32	12 week
ne	29	0-100	62.7	14.3	8.19	3.37	
	31	0-100	60.5	11.3	4.26	3.36	
extrose +	24	0-100	59.9	12.2	14.2	2.7	12 week
rmal ercise)	13	0-100	67.0	10.8	7.0	3.4	
VOMAC P	ain at 1	2-16 week		•			•
	18	0-20	9.5	2.9	-5.0 €	0.78	16 week
	18	0-20	8.7	4.0	-1.9 €	0.73	
	30	0-100	66.8	14.9	11.78	3.62	12 week
ne	29	0-100	66.7	16.1	5.79	3.67	
	31	0-100	63.2	13.1	4.89	3.66	
extrose +	24	0-100	62.3	12.9	15.4	3.0	12 week
rmal ercise)	13	0-100	71.9	13.5	3.5	3.9	
VOMAC F	unction	n at 12–16 we	ek	,			
	18	0-68	33.6	10.7	-14.6 €	2.14	16 week
	18	0-68	26.8	12.8	-3.6 €	2.52	
	30	0-100	65.2	15.8	14.61	3.40	12 week
ne	29	0-100	67.6	17.5	6.63	3.44	
	31	0-100	61.9	12.7	4.89	3.43	
extrose +	24	0-100	62.6	12.9	14.3	2.8	12 week
rmal ercise)	13	0-100	68.5	11.3	7.8	2.8	
ain intens	ity at 10	6–24 week (m	easured on a Vis	ual Analogue	scale VAS of 0–100	mm)	
	18	0-100	48.6	21.8	-29.70 €	4.57	16 week
	18	0-100	38.3	24.8	-9.92 €	4.58	
	36	0-100	21.5	22.4	-5.4	2.4	24 week
	35	0-100	27.3	20.2	-10.4	2.5	
	36	0-100	39.4	28.2	-13.9	3.1	
	35	0-100	38.3	22.0	-9.80	3.2	
	36	0-100	53.3	28.0	-13.7	3.2	
	35	0-100	58.3	26.0	-12.3	3.2	
ain intens	ity at 10		ong Baker/Knee		1		1
	18	WBS 0-5	2.7	1.2	-1.27 €	0.33	16 week
	18	WBS 0-5	2.3	1.1	-0.19 €	0.26	
	43	KPS 0-5	1.8	0.8	-0.92	0.25	24 week
ne	41	KPS 0-5	1.7	0.7	-0.26	0.25	
	47	KPS 0-5	1.7	0.7	-0.26	0.23	
NOM ACC		ite at 52 week		0.0	-0.33	0.24	
VOMACC			1	15.0	15.22	2 22	1
	30	0-100	63.1	15.0	15.32	3.32	
ne	29	0-100	62.7	14.3	7.59	3.36	
extrose ctrose +	24	0-100	59.9	11.3	17.6	3.33	
rmal ercise)	13	0-100	67.0	10.8	8.6	5.0	
VOMAC Pa	ain at 5	2 week					
	30	0-100	66.8	14.9	14.18	3.62	
ne	29	0-100	66.7	16.1	7.38	3.67	
rmal ercise) VOM)	13 AC Pain at 5.	13 0-100 AC Pain at 52 week 30 0-100	13 0-100 67.0 AC Pain at 52 week 30 0-100 66.8	13 0-100 67.0 10.8 AC Pain at 52 week 30 0-100 66.8 14.9	13 0-100 67.0 10.8 8.6 AC Pain at 52 week 30 0-100 66.8 14.9 14.18	13 0-100 67.0 10.8 8.6 5.0 AC Pain at 52 week 30 0-100 66.8 14.9 14.18 3.62

	Outcome: WOMAC Composite at 12–16 week									
	Comparison	n	Scale Range ^	Pre-treat- ment mean score	Standard deviation	Post-treatment adjusted mean change	Standard error	Duration of follow up		
	Exercise	31	0-100	63.2	13.1	9.24	3.63			
Rabago ³⁰	Dextrose (dextrose alone or dextrose + morrhuate)	24	0-100	62.3	12.9	18.1	3.8			
	Control (normal saline or exercise)	13	0-100	71.9	13.5	4.6	5.0			
	Outcome WOMAC f	Outcome WOMAC function at 52 week								
	Dextrose	30	0-100	65.2	15.8	16.25	3.39			
Rabago ²²	Normal saline	29	0-100	67.6	17.5	5.46	3.44			
	Exercise	31	0-100	61.9	12.7	7.31	3.40			
Rabago ³⁰	Dextrose (dextrose alone or dextrose + morrhuate)	24	0-100	62.6	12.9	18.6	2.9			
	Control (normal saline or exercise)	13	0-100	68.5	11.3	9.8	4.8			

Table 4. Effectiveness of prolotherapy injection by comparison type for WOMAC composite, WOMAC pain, WOMAC Function scores and pain intensity. Keys: *The VAS score in Dumais 2012 represents a global assessment of disease activity, with minus sign (−) indicating improvement. *The VAS score in Reeves 2002 represents measurement of pain intensity at rest, with minus sign (−) indicating improvement. *The VAS score in Reeves 2002 represents measurement of pain intensity with walking, with minus sign (−) indicating improvement. *The VAS score in Reeves 2002 represents measurement of pain intensity with stair use, with minus sign (−) indicating improvement. *Wong Baker = Wong Baker Faces Pain Rating scale from 0–5, with minus sign (−) indicating improvement. *The Knee pain scale assessed severity on a 0–5 ordinal scale on each individual knee, with minus sign (−) indicating improvement. *Dumais adopted WOMAC version constructed on a five point Likert scale: composite (0–96), pain (0–20), stiffness (0–8) and function (0–68). Lower score indicates better knee related metrics, minus sign (−) indicates pain reduction. *Rabago adopted WOMAC version constructed on a five point Likert scale: composite (0–96), pain (0–20), stiffness (0–8) and function (0–68); values were then converted to a 0–100 scale for each of the four domains. Composite WOMAC score reflect a weighted average. Higher score indicates better knee related metrics, of which a positive sign (+) indicates improvement. *The values in standard deviation were converted to standard error.

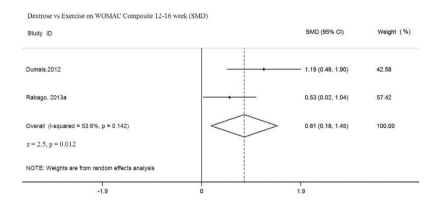


Figure 2. Dextose vs Exercise on WOMAC Composite 12–16 week (SMD).

rigor by increasing the overall sample size, providing a more precise estimate⁶¹. In a narrative review, evaluation of efficacy in RCTs is based primarily on the p-values of individual studies, a so-called 'vote counting' approach, and the risk of arriving at a biased conclusion may even be higher⁶². Because most of the results in the included trials were positive, readers may in the case of narrative review be tempted to draw an over-confident conclusion⁶³. A meta-analysis, with calculation of I² value, is able to express current best evidence and heterogeneity among studies quantitatively⁶⁴. Specifically, we determined that the I² value of WOMAC composite score is 53.6%, indicating considerable heterogeneity on the effect estimate, a finding that can be communicated only through meta-analysis⁶⁵.

Strengths include the conduct of a comprehensive literature search, duplicate study selection and data extraction by two independent reviewers, and appraising risk of bias of all included studies using a validated tool independently by two reviewers.

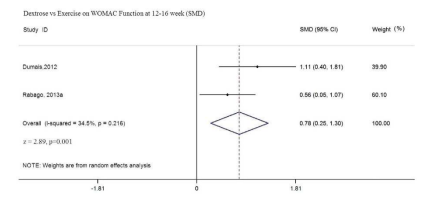


Figure 3. Dextrose vs Exercise on WOMAC Function at 12-16 week (SMD).

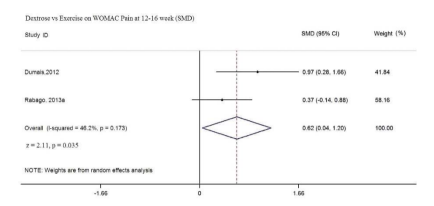


Figure 4. Dextrose vs Exercise on WOMAC Pain at 12–16 week (SMD).

Conclusion

The results of this systematic review indicate that hypertonic dextrose prolotherapy conferred a positive, significant beneficial effect meeting criteria for clinical relevance in the treatment of knee OA, compared with saline injection and exercise. However, moderate heterogeneity existed among these trial results. Larger, long-term trials with uniform outcomes and high methodological standards are needed for more a more comprehensive assessment of the overall treatment effect of prolotherapy.

References

- 1. Brooks, P. M. The burden of musculoskeletal disease-a global perspective. Clin. Rheumatol. 25(6), 778-781(2006).
- 2. Fransen, M. et al. The epidemiology of osteoarthritis in asia. Int. J. Rheum. Dis. 14(2), 113–121 (2011).
- 3. Litwic, A., Edwards, M. H., Dennison, E. M. & Cooper, C. Epidemiology and burden of osteoarthritis. *Br. Med. Bull.* 105, 185–199 (2013).
- 4. Gupta, S., Hawker, G. A., Laporte, A., Croxford, R. & Coyte, P. C. The economic burden of disabling hip and knee osteoarthritis (OA) from the perspective of individuals living with this condition. *Rheumatology (Oxford)*. **44(12)**, 1531–1537 (2005).
- 5. Cross, M. et al. The global burden of hip and knee osteoarthritis: estimates from the global burden of disease 2010 study. Ann. Rheum. Dis. 73(7), 1323–1330 (2014).
- 6. Hochberg, M. C. et al. American college of rheumatology 2012 recommendations for the use of nonpharmacologic and pharmacologic therapies in osteoarthritis of the hand, hip, and knee. Arthrit. Care Res. (Hoboken). 64(4), 465–474 (2012).
- 7. Bruyere, O. et al. An algorithm recommendation for the management of knee osteoarthritis in europe and internationally: a report from a task force of the european society for clinical and economic aspects of osteoporosis and osteoarthritis (ESCEO). Semin. Arthritis. Rheu. 44(3), 253–263 (2014).
- 8. McAlindon, T. E. et al. OARSI guidelines for the non-surgical management of knee osteoarthritis. Osteoarthr. Cartil. 22(3), 363–388 (2014).
- 9. The national academies of sciences, engineering, medicine. Initial national priorities for comparative effectiveness research. (2009) Available at: http://Www.iom.edu/reports/2009/ComparativeEffectivenessResearchPriorities.aspx. (Date of access: April 2014).
- Samson, D. J. et al. Treatment of primary and secondary osteoarthritis of the knee. Evid. Rep. Technol. Assess (Full Rep.). 157, 1–157 (2007).
- 11. Rabago, D., Best, T. M., Beamsley, M. & Patterson, J. A systematic review of prolotherapy for chronic musculoskeletal pain. *Clin. J. Sport Med.* **15(5)**, 376–380 (2005).
- 12. Oh, S. et al. Dextrose-induced subsynovial connective tissue fibrosis in the rabbit carpal tunnel: a potential model to study carpal tunnel syndrome? *Hand (N. Y.).* **3(1)**, 34–40 (2008).
- 13. Yoshii, Y. et al. The effects of hypertonic dextrose injection on connective tissue and nerve conduction through the rabbit carpal tunnel. Arch. Phys. Med. Rehab. 90(2), 333–339 (2009).
- 14. Yoshii, Y. et al. Effects of hypertonic dextrose injections in the rabbit carpal tunnel. J. Orthop. Res. 29(7), 1022-1027 (2011).

- 15. Yoshii, Y. et al. Effects of multiple injections of hypertonic dextrose in the rabbit carpal tunnel: a potential model of carpal tunnel syndrome development. *Hand (N.Y.)*. **9(1)**, 52–57 (2014).
- Park, Y. S. et al. Intra-articular injection of a nutritive mixture solution protects articular cartilage from osteoarthritic progression induced by anterior cruciate ligament transection in mature rabbits: a randomized controlled trial. Arthritis Res. Ther. 9(1), R8 (2007).
- 17. Vora, A., Borg-Stein, J. & Nguyen, R. T. Regenerative injection therapy for osteoarthritis: fundamental concepts and evidence-based review. *P.M. & R.* **4(5, Supplement)**, S104–S109 (2012).
- 18. Rabago, D., Slattengren, A. & Zgierska, A. Prolotherapy in primary care practice. Prim. Care. 37(1), 65-80 (2010).
- 19. Distel, L. M. & Best, T. M. Prolotherapy: A clinical review of its role in treating chronic musculoskeletal pain. P.M. & R. 3(6, Supplement), S78-S81 (2011).
- Jahangiri, A., Moghaddam, F. R. & Najafi, S. Hypertonic dextrose versus corticosteroid local injection for the treatment of
 osteoarthritis in the first carpometacarpal joint: a double-blind randomized clinical trial. J. Orthop. Sci. 19(5), 737–743 (2014).
- Reeves, K. D. & Hassanein, K. Randomized, prospective, placebo-controlled double-blind study of dextrose prolotherapy for osteoarthritic thumb and finger (DIP, PIP, and trapeziometacarpal) joints: evidence of clinical efficacy. J. Altern. Complement. Med. 6(4), 311–320 (2000).
- 22. Rabago, D. *et al.* Hypertonic dextrose and morrhuate sodium injections (prolotherapy) for lateral epicondylosis (tennis elbow): results of a single-blind, pilot-level, randomized controlled trial. *Am. J. Phys. Med. Rehabil.* **92(7)**, 587–596 (2013).
- 23. Topol, G. A. & Reeves, K. D. Regenerative injection of elite athletes with career-altering chronic groin pain who fail conservative treatment: a consecutive case series. *Am. J. Phys. Med. Rehabil.* **87(11)**, 890–902 (2008).
- 24. Kim, W. M., Lee, H. G., Jeong, C. W., Kim, C. M. & Yoon, M. H. A randomized controlled trial of intra-articular prolotherapy versus steroid injection for sacroiliac joint pain. *J. Altern. Complement Med.* **16(12)**, 1285–1290 (2010).
- Dagenais, S., Yelland, M. J., Del Mar, C. & Schoene, M. L. Prolotherapy injections for chronic low-back pain. Cochrane Database Syst. Rev. 2(2), CD004059 (2007).
- 26. Rabago, D. et al. A systematic review of four injection therapies for lateral epicondylosis: prolotherapy, polidocanol, whole blood and platelet-rich plasma. Br. J. Sports Med. 43(7), 471–481 (2009).
- Rabago, D., Mundt, M., Zgierska, A. & Grettie, J. Hypertonic dextrose injection (prolotherapy) for knee osteoarthritis: long term outcomes. Complement Ther. Med. 23(3), 388–395 (2015).
- Dumais, R. et al. Effect of regenerative injection therapy on function and pain in patients with knee osteoarthritis: a randomized crossover study. Pain Med. 13(8), 990–999 (2012).
- Reeves, K. D. & Hassanein, K. Randomized prospective double-blind placebo-controlled study of dextrose prolotherapy for knee osteoarthritis with or without ACL laxity. Altern. Ther. Health Med. 6(2), 68–74 (2000).
- 30. Rabago, D. et al. Dextrose prolotherapy for knee osteoarthritis: a randomized controlled trial. Ann. Fam. Med. 11(3), 229–237
- Rabago, D. et al. Association between disease-specific quality of life and magnetic resonance imaging outcomes in a clinical trial of prolotherapy for knee osteoarthritis. Arch. Phys. Med. Rehabil. 94(11), 2075–2082 (2013).
- 32. Moher, D., Liberati, A., Tetzlaff, J. & Altman, D. G. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. *Int. J. Surg.* 8(5), 336–341(2010).
- 33. Altman, R. D. Criteria for classification of clinical osteoarthritis. J. Rheumatol. Suppl. 27, 10-12 (1991).
- 34. Roos, E. M., Klassbo, M. & Lohmander, L. S. WOMAC osteoarthritis index. reliability, validity, and responsiveness in patients with arthroscopically assessed osteoarthritis, western ontario and MacMaster universities. Scand. I. Rheumatol. 28(4), 210–215 (1999).
- 35. Tubach F. et al. Evaluation of clinically relevant changes in patient reported outcomes in knee and hip osteoarthritis: the minimal clinically important improvement. Ann. Rheum. Dis. 64(1), 29–33 (2005).
- 36. Bellamy, N. et al. Recommendations for a core set of outcome measures for future phase III clinical trials in knee, hip, and hand osteoarthritis. consensus development at OMERACT III. J. Rheumatol. 24(4), 799–802 (1997).
- 37. Bellamy, N., Buchanan, W. W., Goldsmith, C. H., Campbell, J. & Stitt, L. W. Validation study of WOMAC: a health status instrument for measuring clinically important patient relevant outcomes to antirheumatic drug therapy in patients with osteoarthritis of the hip or knee. *J. Rheumatol.* 15(12), 1833–1840 (1988).
- 38. Rejeski, W. J. et al. The evaluation of pain in patients with knee osteoarthritis: the knee pain scale. J. Rheumatol. 22(6), 1124–1129 (1995).
- 39. Wong-Baker Faces. Available at: http://wongbakerfaces.org/. (Date of access: April 2014 (2009)).
- 40. Higgins, J. P. et al. The cochrane collaboration's tool for assessing risk of bias in randomised trials. B.M.J. 343, 5928 (2011).
- 41. Wallace, B. C., Schmid, C. H., Lau. J. & Trikalinos, T. A. Meta-analyst: software for meta-analysis of binary, continuous and diagnostic data. B.M.C. Med. Res. Methodol. 9(1), 80 (2009).
- 42. Borenstein, M., Hedges, L. V., Higgins, J. P. T. & Rothstein, H. R. A basic introduction to fixed-effect and random-effects models for meta-analysis. *Res Synth Methods*. 1(2), 97–111 (2010).
- 43. Deeks, J. J., Higgins, J. P. T. & Altman, D. G. Heterogeneity in *Cochrane handbook for systematic reviews of interventions*. (eds Higgins, J. P. T. & Green, S.) Ch.9.5, 276–282 (Wiley-Blackwell, 2008).
- 44. Tubach, F., Wells, G. A., Ravaud, P. & Dougados, M. Minimal clinically important difference, low disease activity state, and patient acceptable symptom state: methodological issues. *J. Rheumatol.* 32(10), 2025–2029 (2005).
- 45. Ehrich, E. W. et al. Minimal perceptible clinical improvement with the western ontario and McMaster universities osteoarthritis index questionnaire and global assessments in patients with osteoarthritis. *J. Rheumatol.* 27(11), 2635–2641 (2000).
- 46. Brandt, K. D., Fife, R. S., Braunstein, E. M. & Katz, B. Radiographic grading of the severity of knee osteoarthritis: relation of the kellgren and lawrence grade to a grade based on joint space narrowing, and correlation with arthroscopic evidence of articular cartilage degeneration. *Arthritis Rheum.* 34(11), 1381–1386 (1991).
- 47. Rabago, D. et al. Hypertonic dextrose injections (prolotherapy) for knee osteoarthritis: results of a single-arm uncontrolled study with 1-year follow-up. J Altern. Complement Med. 18(4), 408–414 (2012).
- 48. Faraone, S. V. Interpreting estimates of treatment effects; implications for managed care. P. & T. 33(12), 700 (2008).
- Borenstein, M., Hedges, L. V., Higgins, J. P. T. & Rothstein, H. R. Identifying and quantifying heterogeneity in *Introduction to meta-analysis*. Ch. 16, 107–125. (John Wiley & Sons, Ltd, 2009).
- 50. Noordzij, M. et al. Sample size calculations: basic principles and common pitfalls. Nephrol. Dial. Transplant. 25(5), 1388–1393 (2010).
- 51. Banks, A. A rationale for prolotherpay. J. Orthop. Med. 13(3), 54-59 (1991).
- 52. Vora, A., Borg-Stein, J. & Nguyen. R. T. Regenerative injection therapy for osteoarthritis: fundamental concepts and evidence-based review. P.M. & R. 4(5, Supplement), S104–S109 (2012).
- 53. Dobson, F. et al. OARSI recommended performance-based tests to assess physical function in people diagnosed with hip or knee osteoarthritis. Osteoarthr. Cartil. 21(8), 1042–1052 (2013).
- 54. Hoogeboom, T. J. et al. Joint-pain comorbidity, health status, and medication use in hip and knee osteoarthritis: a cross-sectional study. Arthrit. Care Res. 64(1), 54–58 (2012).
- 55. Hośnijeh, F. S., Runhaar, J., van Meurs, J. B. & Bierma-Zeinstra, S. M. Biomarkers for osteoarthritis: can they be used for risk assessment? a systematic review. *Maturitas.* 82(1), 36–49 (2015).

- 56. Xie, F. et al. Use of a disease-specific instrument in economic evaluations: mapping WOMAC onto the EQ-5D utility index. Value Health. 13(8), 873–878 (2010).
- 57. Jancsó, G. Neurogenic mechanisms in arthritis in *Neurogenic inflammation in health and disease* (eds Donaldson, L. F.) Ch.4, 211–241 (Elsevier, 2009).
- 58. Lyftogt, J. Subcutaneous prolotherapy treatment of refractory knee, shoulder and lateral elbow pain. *Australas Musculoskelet. Med.* 12(2), 110–112 (2007).
- 59. Qvistgaard, E. et al. Guidance by ultrasound of intra-articular injections in the knee and hip joints. Osteoarthr. Cartil. 9(6), 512–517 (2001)
- 60. Guyatt, G. H. et al. GRADE guidelines 6. rating the quality of evidence-imprecision. J. Clin. Epidemiol. 64(12), 1283-1293 (2011).
- 61. Deeks, J. J., Higgins, J. P. T. & Altman, D. G. Analysing data and undertaking meta-analyses in *Cochrane handbook for systematic reviews of interventions*. (eds Higgins, J. P. T. & Green, S.) Ch.9, 243–296 (Wiley-Blackwell, 2008).
- 62. Ioannidis, J. P., Patsopoulos, N. A. & Rothstein, H. R. Reasons or excuses for avoiding meta-analysis in forest plots. B.M.J. 336(7658), 1413–1415 (2008)
- 63. Borenstein, M., Hedges, L. V., Higgins, P. T. & Rothstein, H. R. Vote Counting–A new name for an old problem in *Introduction to meta-analysis*. 251–255 (John Wiley & Sons, 2009).
- 64. Borenstein, M., Hedges, L. V., Higgins, P. T. & Rothstein, H. R. Criticisms of meta-analysis in *Introduction to meta-analysis*.377–387 (John Wiley & Sons, 2009).
- 65. Deeks, J. J., Higgins, J. P. T. & Altman, D. G. Identifying and measuring heterogeneity in Cochrane handbook for systematic reviews of interventions. (eds Higgins, J. P. T. & Green, S.) Ch. 9.5.2, 277–278 (Wiley-Blackwell, 2008).

Author Contributions

R.W.S.S. is the first author of this article, responsible for the study design, review of literatures and completion of manuscript. V.C.H.C. is the corresponding author in this study; he provides methodology support and acts as another reviewer in literatures search K.D.R. and D.R. are the authors of the three papers involved in this systematic review; both of them provide valuable advice on the study design, data interpretation and discussion. K.K.W.C. provides expert opinions on the practical aspect of prolotherapy and S.Y.S.W. as a senior author to overlook the whole study. D.C.C.C., R.S.T.H. and X.W. provide full statistical support in this article.

Additional Information

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OPEN Erratum: Hypertonic dextrose injections (prolotherapy) in the treatment of symptomatic knee osteoarthritis: A systematic review and meta-analysis

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This Article contains errors in Tables 1, 2, 3 and 4 where references 29, 30 and 31 are incorrectly given as references 21, 22 and 30 respectively.

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