

# Do knee abnormalities visualised on MRI explain knee pain in knee osteoarthritis? A systematic review

Erlangga Yusuf,<sup>1</sup> Marion C Kortekaas,<sup>1</sup> Iain Watt,<sup>2</sup> Tom W J Huizinga,<sup>1</sup> Margreet Kloppenburg<sup>1</sup>

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<sup>1</sup>Department of Rheumatology, Leiden University Medical Center, Leiden, The Netherlands  
<sup>2</sup>Department of Radiology, Leiden University Medical Center, Leiden, The Netherlands

## Correspondence to

Erlangga Yusuf, Department of Rheumatology, Leiden University Medical Center, C1-46, Postbus 9600, 2300 RC Leiden, The Netherlands; [e.yusuf@lumc.nl](mailto:e.yusuf@lumc.nl)

EY and MCK contributed equally to this work.

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## ABSTRACT

**Objective** To systematically evaluate the association between MRI findings (cartilage defects, bone marrow lesions (BML), osteophytes, meniscal lesion, effusion/synovitis, ligamentous abnormalities, subchondral cysts and bone attrition) and pain in patients with knee osteoarthritis (OA) in order to establish the relevance of such findings when assessing an individual patient.

**Methods** The Medline, Web of Science, Embase and Cumulative Index to Nursing & Allied Health Literature (CINAHL) databases up to March 2010 were searched without language restriction to find publications with data on the association between MRI findings of knee OA (exposure of interest) and knee pain (outcome). The quality of included papers was scored using a predefined criteria set. The levels of evidence were determined qualitatively using best evidence synthesis (based on guidelines on systematic review from the Cochrane Collaboration Back Review Group). Five levels of evidence were used: strong, moderate, limited, conflicting and no evidence.

**Results** A total of 22 papers were included; 5 had longitudinal and 17 cross-sectional data. In all, 13 reported a single MRI finding and 9 multiple MRI findings. Moderate levels of evidence were found for BML and effusion/synovitis. The OR for BML ranged from 2.0 (no CI was given) to 5.0 (2.4 to 10.5). The OR of having pain when effusion/synovitis was present ranged between 3.2 (1.04 to 5.3) and 10.0 (1.1 to 149). The level of evidences between other MRI findings and pain were limited or conflicting.

**Conclusions** Knee pain in OA is associated with BML and effusion/synovitis suggesting that these features may indicate the origin of pain in knee OA. However, due to the moderate level of evidence these features need to be explored further.

## INTRODUCTION

Knee is the major site of osteoarthritis (OA), the most common rheumatic disorder which is characterised by pain that leads to significant restriction in patients' daily activity.<sup>1 2</sup> Despite its importance, the source of pain remains unclear.<sup>3</sup> To treat OA optimally, knowledge of the source of pain is important since new therapies can be specifically targeted.

An important element in understanding pain is to know which structures produce it inside the knee since the pathology of knee OA involves the whole knee joint.<sup>3</sup> To assess knee structures in vivo imaging modalities are needed. On radiographs, hallmarks of knee OA such as bony outgrowth and cartilage loss, which are visualised as osteophytes and joint space narrowing, respectively, do

not show a consistent association with knee pain.<sup>4</sup> Other potential sources include abnormalities in subchondral bone, ligamentous damage, meniscal injury and synovitis.<sup>5</sup> However, these potential sources cannot be assessed on conventional radiographs. More advanced imaging techniques are needed currently best exemplified by MRI.

Several studies have investigated MRI findings related to pain but to our knowledge, no summarisation of data has been performed in a systematic manner. Such a review requires a focused research question, an explicit research strategy and a system to evaluate the quality of evidence.<sup>6</sup> Therefore, we sought to evaluate the relationship between MRI findings in knee OA and knee pain. We summarised eight commonly reported MRI findings: cartilage defects, bone marrow lesions (BML), osteophytes, meniscal lesion, effusion/synovitis, ligamentous abnormalities, subchondral cysts and bone attrition (see supplementary material).

## MATERIALS AND METHODS

The present review is systematic review of observational studies. Therefore, we adhered to a protocol developed from a widely recommended method for systematic review/meta-analysis of observational studies (MOOSE).<sup>7</sup> We included studies with data on the association between MRI features of knee OA (exposure of interest) and knee pain (outcome). The following studies were excluded: reviews, abstracts, letters to the editor, case reports, case series and studies concerning study population with other underlying musculoskeletal diseases.

## Data sources, searches and extraction

Using the following key words: 'knee', 'knee pain', 'MRI', 'osteoarthritis' in combination with all possible key words concerning MRI features we wanted to investigate, we searched the following medical databases up to March 2010: Medline (from 1966), Science Citation Index through Web of Science (from 1945), Embase (from 1980) and, Cumulative Index to Nursing & Allied Health Literature (CINAHL) (from 1982). No language restriction was applied and no search of unpublished studies was performed. Additionally, the reference lists of all relevant identified articles were screened and Google Scholar was searched to find additional papers. Complete search strategies can be obtained from the authors on request.

Two reviewers, EY (a PhD student) and MCK (a rheumatologist) independently screened the titles of retrieved references for obvious exclusion and read the remaining abstract to determine eligible

studies. Differences were solved by discussion or by consulting a third reviewer (MK, a senior rheumatologist).

From eligible papers, information was collected on the following categories: (i) type of study, performed by looking at the method of data analysis (when a study provided data on the association between MRI features change in time with change in pain level in time, the study was considered to be a prospective cohort study; if this analysis was not available, such as in a case-control study, the study was regarded to be of a cross-sectional design); (ii) study population (patient characteristics, size, gender and age); (iii) definition of knee OA; (iv) assessment of MRI findings; (v) assessment of pain; (vi) potential confounders; and (vii) results of the association between MRI features and pain.

### Assessment of study quality

Independently, the same two reviewers assessed the methodological quality of included studies using a predefined criteria set which was previously used in systematic reviews in the area of musculoskeletal disorders (see supplementary material).<sup>8 9</sup> Several domains were assessed: population, selection bias, assessment of determinants on MRI, assessment of the outcome, follow-up analysis and data presentation.

For each criterion met in the article, a '1' was given; otherwise, a '0' was given. We defined rules on how to assess specific situations. A study could describe multiple MRI features but not all were assessed reproducibly (criterion 5) or using standardised criteria (criterion 6). For such a study, the criteria are scored as a proportion of MRI features which were assessed reproducibly or using standardised criteria from the total MRI features investigated.

Differences in scoring were resolved by discussion or by consulting the third reviewer. Maximum scores possible were 11 for prospective cohort and 9 for cross-sectional study design. The total score for a study (in %) is the total score given for a study divided by the maximum possible score. The mean of the quality scores of all studies, which was 62%, was used to classify studies as high or low quality.

### Rating the body of evidence

The summary of evidence for each MRI feature was given by using best evidence synthesis based on the guidelines on systematic review of the Cochrane Collaboration Back Review Group.<sup>10</sup> This is an alternative to pooling of association sizes when the included studies were heterogenous.<sup>8</sup> The synthesis has five levels of evidence: (1) strong, when general consistent findings were reported in multiple high-quality cohort studies; (2) moderate, when one high-quality cohort study and at least two high-quality cross-sectional studies show general consistent findings or when at least three high quality cross-sectional studies who general consistent findings; (3) limited, when general consistent findings were found in a single cohort study, or in maximum two cross-sectional studies; (4) conflicting, when no consistent findings were reported; and (5) no evidence, when no study could be found. This synthesis puts more weight on a prospective cohort design which is appropriate for our review question since it takes into account the change in determinant (MRI feature) and change in outcome (pain).

Sensitivity analyses by defining other cut-offs (median score of all studies instead of mean) of high quality studies were performed. We also present the number of positive studies without quality assessment to give readers the opportunity to compare this with the best evidence synthesis results.

A study that investigated multiple features was counted as a single study for each MRI feature investigated. A study was regarded as positive if it showed a significant association between an MRI feature and knee pain. When a study included subfeatures of an MRI finding, that is, tear and subluxation for meniscal lesion, the study was regarded as positive when at least one of these showed positive association. Since effusion and synovitis cannot be readily differentiated on non-enhanced MRI,<sup>9 11</sup> we analysed these features together.

## RESULTS

### Literature flow

After screening their title, 2144 of 2629 identified references were excluded (figure 1). From the 485 remaining references, 19 papers were included. We selected the most recent publication<sup>12</sup> of two publications with overlapping results.<sup>12 13</sup> Four publications<sup>14-17</sup> came from the same authors and used the same patient population. We therefore selected two of them.<sup>14 16</sup> These two selected studies defined cartilage loss as determinant and pain as outcomes, contradictory to the two others which defined the determinant and outcomes conversely. After additional searching, another three papers were found.<sup>16 18 19</sup> In total, 22 papers were selected. In all, 5 studies reported longitudinal data<sup>12 14 16 20 21</sup> and 17<sup>18 19 22-36</sup> were cross-sectional studies.

### Characteristics of included studies

Of the 22 analysed papers, 8 published associations of multiple MRI features (table 1),<sup>19 25 26 29 30 32 34 36</sup> the others investigated only a single MRI feature.

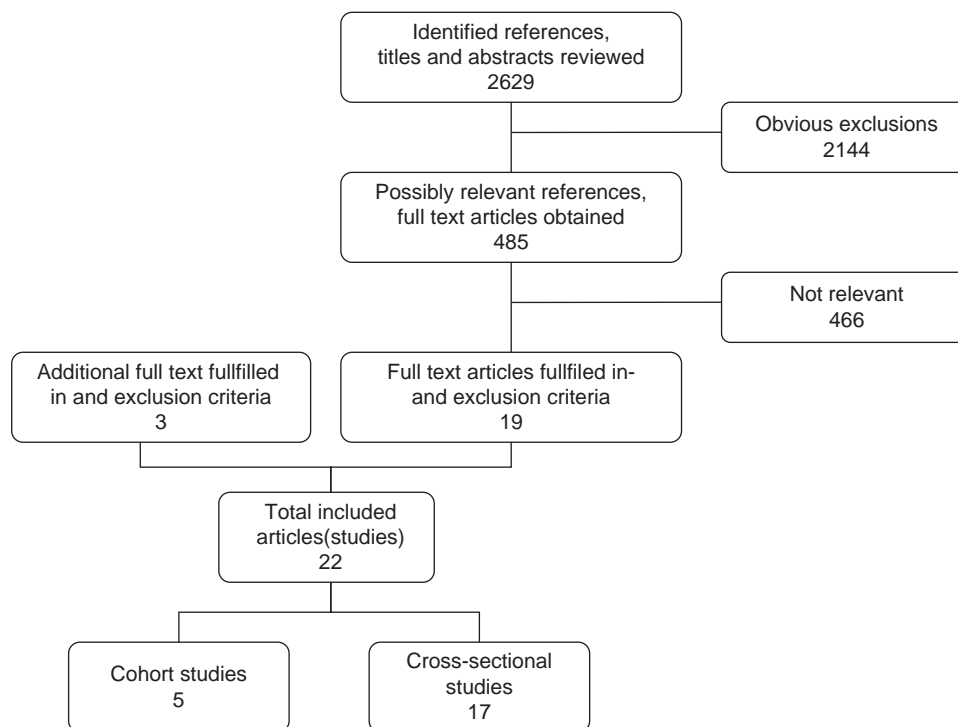
Of these papers (table 1), 10 were results from 3 studies: the Boston Osteoarthritis Knee Study (BOKS),<sup>12 18 22 24 28 33</sup> the Southeast Michigan OA (SEM) cohort,<sup>26 34</sup> and the Genetic Arthrosis Progression Study (GARP).<sup>20 29</sup> Most studies used a General Electric MRI system (in 14 publications.<sup>12 13 16 18 19 22-24 26 28 30 32-34</sup> A Siemens MRI system was used in six publications<sup>14 25 27 31</sup> and a Philips MRI system was used in two publications.<sup>20 29</sup> Two studies<sup>35 36</sup> used a 3 T magnetic field system, all others used a 1.5 T system. Only one study<sup>35</sup> used MRI contrast agent.

Patients investigated in the included studies were of both sexes and older than 50 years, except for one which studied women alone with mean age of 47 years (table 1).<sup>26</sup> Almost all studies defined knee OA by using clinical and radiographic criteria of American College of Rheumatology, which requires at least knee pain and osteophyte on radiograph. Only five studies defined knee OA purely radiographically.<sup>19 23 26 27 31</sup>

### Study quality assessment

We agreed on 212 of 227 (93%) quality assessment items scored (see supplementary material). Most disagreement focused on the clarity of description of the study population (criterion 2) and participation rate (criterion 3).

In general, many publications either did not assess MRI findings using standardised and validated criteria or they did not inform the reader about this (criterion 5). In many prospective cohort studies the researchers were not blinded for the time order of MRI scans (criterion 7) and differences between withdrawal and completed groups were not described (criterion 10). In cross-sectional studies, the most common limitations were participation rate (criterion 3) and lack of adjustment of possible confounders such as age and sex (criterion 11).



**Figure 1** Results of literature research

### Association between MRI features and pain (best-evidence synthesis)

#### Cartilage defect

Six studies<sup>19 26 29–32</sup> investigated cartilage defects using semiquantitative scores, five<sup>14 16 23 25 34</sup> used quantitative methods and one used quantitative method on contrast-enhanced MRI.<sup>35</sup> The level of evidence on the association between cartilage defects and pain was conflicting: three<sup>16 19 34</sup> of five high-quality studies showed a positive association with pain. When all 12 studies which investigated cartilage defects<sup>14 16 19 23 25 27 29–32 34 35</sup> were summarised, 50% showed a positive association independent of study quality.

#### Bone marrow lesions

The evidence about the association between BML and pain was moderate. Four<sup>19 24 34 36</sup> of five high-quality studies showed an association between BML and pain. One high-quality cohort study showed no association.<sup>20</sup> Three of the four high-quality cross-sectional studies that demonstrated a positive association presenting an OR as quantitative measure of association. The OR ranged from 2.0 (adjusted for effusion and synovitis)<sup>36</sup> to 5.0 (unadjusted, 95% CI 2.4 to 10.5).<sup>34</sup> One study reported a  $\beta$  coefficient of 3.72 (95% CI 1.76 to 5.68).<sup>19</sup> When all eight studies investigating BML<sup>19 20 24 26 30 32 34 36</sup> were taken into account 63% reported a positive association between BML and pain.

#### Osteophytes

Neither of the two high-quality studies showed a positive association between osteophytes with pain.<sup>29 33</sup> According to best evidence synthesis this gives limited level of evidence on the no association between osteophytes and knee pain.

#### Meniscal lesions

Only one<sup>19</sup> of three high-quality cross-sectional studies showed a positive association resulting in a conflicting level of evidence for the association between meniscal lesions and pain.<sup>18 19 29</sup>

When all studies were taken into account; 33% showed a positive association.

#### Synovitis/joint effusion

A moderate association was found for effusion/synovitis, since all four<sup>12 19 29 36</sup> high-quality studies showed a positive association. One of which was a high-quality cohort study.<sup>12 19 29</sup> This study performed separate analyses for effusion and synovitis: the analysis between effusion and pain showed no association whereas the association between synovitis and pain was positive. We regarded this study as positive, because we deemed a study as a positive study when at least one of the subfeatures showed a positive association. Four high-quality studies reported quantitative measures of association. Three reported the OR of having pain when effusion/synovitis was present, ranging between 2.6 (adjusted for synovitis and BML)<sup>36</sup> and 10.0 (adjusted for age, sex BMI and intrafamily effects, 99% CI 1.13 to 149).<sup>29</sup> One other study reported  $\beta$  regression of 9.82 (95% CI 0.38 to 19.27).<sup>19</sup> When no quality assessment was performed, 86% of included studies<sup>12 19 21 25 26 29 30 36</sup> showed a positive association with pain.

#### Ligament disease

Two studies<sup>28 30</sup> classified ligament abnormalities as presence or absence of tears, and three studies<sup>19 22 26</sup> used semiquantitative scores. Since only two high-quality studies<sup>19 22</sup> were available, which showed positive association, this resulted in a limited level of evidence for a positive association between ligament abnormalities and pain. When all five studies<sup>19 22 26 30</sup> were taken in account, only 40% showed a positive association.

#### Subchondral cyst

Subchondral cysts were not associated with pain. Two high-quality studies showed no association and this resulted in a limited level of evidence.<sup>19 29</sup>

**Table 1** Characteristics of included studies (listed alphabetically by first author surname)

Studies	Features assessed	Pain assessment	Statistical analysis	Quality score (%)	
<b>Cohort studies</b>					
Hill <i>et al</i> <sup>12</sup>	Patients with knee OA (ACR criteria). n=270 (42% women); mean age 67±9 years. BOKS, USA.	Effusion/synovitis	VAS	Linear regression	68
Kornaat <i>et al</i> <sup>20</sup>	Generalised patients with OA. n=182 (86% women); median age 60 years (range: 43–77). GARP study, The Netherlands.	BML	WOMAC pain	Linear mixed model	64
Pelletier <i>et al</i> <sup>21</sup>	Patients with knee OA (ACR criteria) from outpatient rheumatology clinic, n=27 (52% women), mean age 64±9.6 years. Canada.	Synovitis	WOMAC and VAS pain	Spearman correlation	36
Raynauld <i>et al</i> <sup>14</sup>	Patients with knee OA (ACR criteria). n=40 (88% women); mean age 62±8 years. Canada.	Cartilage	WOMAC and VAS pain	Spearman correlation	64
Wluka <i>et al</i> <sup>16</sup>	Patients with knee OA (ACR criteria). n=132 (54% women); mean age 63 years (range 41–86) Australia.	Cartilage	WOMAC pain	Spearman correlation	64
<b>Cross-sectional studies</b>					
Anandacoomarasamy <i>et al</i> <sup>25</sup>	Obese patients with knee OA from general population (ACR criteria), n=77 (68% women), mean age: 51±12.7 years, Sydney, Australia.	Cartilage	WOMAC pain	Spearman correlation	67
Amin <i>et al</i> <sup>22</sup>	BOKS, USA. See above. n=265 (43% women); mean age 67±9 years.	ACL tear	VAS	Student t test	67
Bhattacharyya <i>et al</i> <sup>18</sup>	Cases: BOKS, USA. See above. n=154, mean age: 65 years. Controls: no knee pain, n=49 mean age: 67 years.	Meniscal tear	VAS	Student t test	67
Dunn <i>et al</i> <sup>23</sup>	Patients suspected for clinical OA. n=55 (55% women); mean age 63±3 years. USA.	Cartilage	WOMAC pain	Spearman correlation	22
Felson <i>et al</i> <sup>24</sup>	BOKS, USA. See above. n=401 (33% women in knee pain group, 48% in no pain group); mean age: 62 years (range: 22–91).	BML	Presence/absence of pain	Logistic regression	75
Fernandez-Madrid <i>et al</i> <sup>25</sup>	Case: patients with knee OA (ACR criteria). n=52 (67% women); mean age 55±14 years. Control: general population. n=40 (62% women), 49±15 years. Detroit, USA.	Cartilage, osteophytes, subchondral lesions, effusion/synovitis, meniscal tears	Presence/absence of pain	χ <sup>2</sup> test	72
Hayes <i>et al</i> <sup>26</sup>	Four groups (each n=30, 100% women): no pain, no radiographic knee OA, mean age 45±1 years; no pain, radiographic knee OA, 46±1 years; pain, no radiographic knee OA, 47±1 years; pain, radiographic knee OA, 47±1 years. Southeast Michigan Osteoarthritis cohort, USA.	Cartilage, osteophytes, subchondral cysts, BML, effusion/synovitis, meniscal tear, ACL tear	Presence/absence of pain	Fisher exact test of general association	56
Hernández-Molina <i>et al</i> <sup>27</sup>	Patients with knee OA (K&L ≥2). n=1273 (48% women); mean age: 65±9 years. Framingham OA study cohort, Massachusetts, USA.	Bone attrition	Presence/absence of pain	χ <sup>2</sup> test	78
Hill <i>et al</i> <sup>28</sup>	Cases: BOKS, USA. See above. n=360, 33% women, mean age: 68 years. Controls: no knee pain. n=73, 65% with K&L ≥2 and JSN≥1, 57% men, 66 years.	ACL tear	Presence/absence of pain	χ <sup>2</sup> test	50
Kornaat <i>et al</i> <sup>29</sup>	GARP. See above. n=205 (80% women); median age 60 years (range: 43–77).	Cartilage, osteophytes, subchondral cysts, BML, effusion, meniscal defects	Presence/absence of pain	Logistic regression	78
Link <i>et al</i> <sup>30</sup>	Patients with knee OA (ACR criteria). n=50 (60% women); mean age 64±11 years.	Cartilage, BML, meniscal tear, ACL tear	WOMAC pain	Wilcoxon rank sum test	47
Lo <i>et al</i> <sup>36</sup>	Patients with knee OA (Knee pain or stiffness and osteophytes OARS atlas score 1–3), n=160 (50% women), mean age 61±9.9. OA initiative.	BML, effusion/synovitis	WOMAC pain	Logistic regression	78
Pelletier <i>et al</i> <sup>31</sup>	Knee OA (radiographic) from general population. Subset from clinical trial on Risendronate in North America. n=110 (64% women); mean age 62±7 years.	Cartilage	WOMAC pain	Spearman correlation	39
Phan <i>et al</i> <sup>32</sup>	Patients with knee OA (ACR criteria), n=34 and general population, n=6, 60% women, mean age: 58±16 years.	Cartilage, BML	WOMAC pain	Correlation not specified	67
Sengupta <i>et al</i> <sup>33</sup>	BOKS. See above. n=217 (30% women); mean age 67±9 years.	Osteophytes	10-point pain scale	Logistic regression	78
Sowers <i>et al</i> <sup>34</sup>	Southeast Michigan Osteoarthritis cohort, USA. See above.	Cartilage, BML	VAS pain	Wilcoxon or Maentel–Haenszel test of general association	78
Torres <i>et al</i> <sup>19</sup>	Patients with knee OA (K&L ≥2 and ‘a little difficulty’ in one or two WOMAC physical function scale). n=143 (88% women); mean age 70±10 years.	Cartilage, osteophytes, bone cysts, bone attrition, BML, synovitis, meniscal tears, ligament abnormalities (MCL, LCL and ACL)	VAS pain	Median quantile regression	78

ACR clinical and radiographic criteria requires knee pain and osteophytes on radiograph.<sup>50</sup>

ACL, anterior cruciate ligament; ACR, American College of Rheumatology; BMI, body mass index; BML, bone marrow lesion; BOKS, Boston OA of the knee study; GARP, Genetic Arthrosis Progression Study; JSN, joint space narrowing; K&L, Kellgren and Lawrence Osteoarthritis Scoring System for knee radiographs; LCL, lateral cruciate ligament; MCL, medial cruciate ligament; n, number of study population; OA, osteoarthritis; VAS, visual analogue scale; WOMAC, Western Ontario and McMaster University.

**Table 2** Best evidence synthesis (MRI features arranged from top to bottom according to the number of studies included)

Studies	Study design	Association (sizes)			Number of studies: positive/total (%)	
		Crude	Adjusted	Adjusted confounders	All	High quality
<b>Cartilage defects (level of evidence: conflicting)</b>						
Scored using semiquantitative scores						
Pelletier <i>et al</i> <sup>31</sup>	CS	r=0.09, p=0.38	–	NA	6/12 (50%)	3 (1 C, 2 CS)/6 (2 C, 3 CS) (50%)
Phan <i>et al</i> <sup>32</sup>	CS	r Not mentioned, NS	–	NA		
<b>Torres <i>et al</i><sup>19</sup></b>	CS	β=1.03 (95% CI 0.6 to 1.5)	0.53 (0.08 to 0.98)	Age and BMI		
Hayes <i>et al</i> <sup>26</sup>	CS	Positive, p=0.001	–	NA		
<b>Kornaat <i>et al</i><sup>29</sup></b>	CS	–	OR 1.12 (99% CI 0.4 to 3.2)	Age, sex, BMI, intrafamily effects		
Link <i>et al</i> <sup>30</sup>	CS	positive, p<0.05	–	NA		
Scored quantitatively						
<b>Raynaud <i>et al</i><sup>14</sup></b>	C	r=–0.25, NS (WOMAC), r=0.12, NS (VAS)	–	NA		
<b>Wluka <i>et al</i><sup>16</sup></b>	C	r=0.28, positive, p=0.002	–	NA		
Fernandez-Madrid <i>et al</i> <sup>25</sup>	CS	NS	–	NA		
<b>Sowers <i>et al</i><sup>34</sup></b>	CS	Positive, p<0.0001	–	NA		
Dunn <i>et al</i> <sup>23</sup>	CS	Positive, p<0.05	–	NA		
Scored using other methods (ie, quantitatively after giving contrast agent)						
<b>Anadacoomarasamy <i>et al</i><sup>35</sup></b>	CS	r=–0.21, p=0.07	–	NA		
<b>Bone marrow lesion (level of evidence: moderate)</b>						
<b>Kornaat <i>et al</i><sup>20</sup></b>	C	–	Mean difference (increasing BML)=2 (95% CI –8 to 11)	Age, sex, BMI, intrafamily effects	5/8 (63%)	4 (CS)/5 (1 C, 4 CS) (80%)
Hayes <i>et al</i> <sup>26</sup>	CS	Positive, p=0.001	–	NA		
<b>Felson <i>et al</i><sup>24</sup></b>	CS	–	OR=3.31 (95% CI 1.5 to 7.4)	Age, sex, radiological and effusion score		
Link <i>et al</i> <sup>30</sup>	CS	p>0.05	–	NA		
<b>Lo <i>et al</i><sup>36</sup></b>	CS	Positive, RR BML scores versus no BML: 1:1.3 2:2.1 3:2.3 p For trend=0.0009	Positive: 1:1.2 2:1.9 3:2.0 p For trend 0.006	Effusion and synovitis		
Phan <i>et al</i> <sup>32</sup>	CS	r Not mentioned, NS	–	NA		
<b>Sowers <i>et al</i><sup>34</sup></b>	CS+	OR=5.0 (95% CI 2.4 to 10.5)	–	NA		
<b>Torres <i>et al</i><sup>19</sup></b>	CS+	β=5.0 (95% CI 3.0 to 7.0)	3.72 (1.8 to 5.7)	Age and BMI		
<b>Osteophytes (level of evidence: limited)</b>						
Presence						
Fernandez-Madrid <i>et al</i> <sup>25</sup>	CS	NS	–	NA	2/6 (33%)	0/2 (CS) (0%)
Hayes <i>et al</i> <sup>26</sup>	CS	Positive, p<0.001	–	NA		
<b>Kornaat <i>et al</i><sup>29</sup></b>	CS	–	OR=1.05 (99% CI 0.4 to 2.9)	Age, sex, BMI, intrafamily effects		
Link <i>et al</i> <sup>30</sup>	CS	p>0.05	–	NA		
Torres <i>et al</i> <sup>19</sup>	CS	β=1.2 (95% CI 0.6 to 1.7)	B=0.5 (0.07 to 0.94)	Age and BMI		
Signal strength						
<b>Sengupta <i>et al</i><sup>33</sup></b>	CS	–	PR=0.94 (0.8 to 1.1)	Age, sex, BMI		
<b>Meniscal lesion (level of evidence: conflicting)</b>						
<b>Bhattacharyya <i>et al</i><sup>18</sup></b>	CS	–	p=0.7	Age	2/6 (33%)	1/3 (CS) (33%)
Fernandez-Madrid <i>et al</i> <sup>25</sup>	CS	NS	–	NA		
Hayes <i>et al</i> <sup>26</sup>	CS	Positive, p=0.001	–	NA		
<b>Kornaat <i>et al</i><sup>29</sup></b>	CS	–	Tears: OR=1.26 (99% CI 0.6 to 2.7), subluxation: OR=1.03 (99% CI 0.5 to 2.2)	Age, sex, BMI, intrafamily effects		
Link <i>et al</i> <sup>30</sup>	CS	p>0.05	–	NA		
<b>Torres <i>et al</i><sup>19</sup></b>	CS	Tears: β=3.3 (95% CI 0.9 to 5.8) Subluxation: β=15.0 (95% CI –0.3 to 30.3)	Tears: 2.0 (0.6 to 3.4) Subluxation: 2.22 (–6.9 to 11.3)	Age and BMI		
<b>Effusion and synovitis (level of evidence: moderate)</b>						
<b>Hill <i>et al</i><sup>12</sup></b>	C	–	Effusion: OR=1.2 (95% CI –8.1 to 10.5), synovitis: OR=3.2 (95% CI 1.04 to 5.3)	Age, sex, BMI, cartilage score at baseline, effusion score, BML score, change in effusion and BML score.	6/8 (80%)	4 (1 C, 3 CS)/4 (1 C, 3 CS) (100%)
Fernandez-Madrid <i>et al</i> <sup>25</sup>	CS	Effusion: positive, p<0.001 Synovitis: NS	–	NA		
Hayes <i>et al</i> <sup>26</sup>	CS	Effusion: positive, p<0.001 Synovitis: positive, p<0.001	–	NA		

Continued

Table 2 continued

Studies	Study design	Association (sizes)			Number of studies: positive/total (%)	
		Crude	Adjusted	Adjusted confounders	All	High quality
<b>Kornaat et al<sup>29</sup></b>	CS	–	Effusion: OR=10.0 (99% CI 1.1 to 149)	Age, sex, BMI, intrafamily effects		
Link et al <sup>27</sup>	CS	Effusion: p>0.05	–	NA		
<b>Lo et al<sup>36</sup></b>	CS+	Effusion: RR BML scores versus no BML: 1:1.8 2:2.4 3:3.1 p For trend<0.0001 Synovitis: 1:1.9 2:1.9 3:2.3 p For trend 0.20	1:1.7 2:2.0 3:2.6 p For trend=0.0004 Synovitis: 1:1.4 2:1.5 3:1.9 p For trend=0.22	Age and BMI		
<b>Torres et al<sup>19</sup></b>	CS	β=15.0 (95% CI –8.2 to 38.2)	9.8 (0.4 to 19.3)	Age and BMI		
Pelletier et al <sup>21</sup>	C	Effusion: r=0.07, positive, p=0.71 (WOMAC); r=0.01, positive, p=0.93 (VAS)	–	NA		
Knee ligament abnormalities (level of evidence: limited)						
<b>Amin et al<sup>22</sup></b>	CS	–	ACL: positive, p<0.05	Age, sex, BMI and cartilage scores	2/5 (40%)	2/2 (CS) (100%)
Hill et al <sup>28</sup>	CS	ACL: positive, p=0.0004	–	NA		
Link et al <sup>30</sup>	CS	ACL: p>0.05, MCL and LCL: p>0.05	–	NA		
<b>Torres et al<sup>19</sup></b>	CS	β (95% CI) ACL: 5.0 (–13.0 to 23.0) MCL: 0 (–11.9 to 11.9) LCL: 15.0 (95% CI –8.2 to 38.2)	ACL: 6.8 (–5.4 to 19.0) MCL: –6.10 (–14.0 to 1.7) LCL: 29.5 (17.8 to 41.1)	Age and BMI		
Hayes et al <sup>26</sup>	CS	ACL and PCL: p=0.23, MCL and LCL, p=0.86	–	NA		
Subchondral cysts (level of evidence: limited)						
Hayes et al <sup>26</sup>	CS	Positive, p<0.001	–	NA	1/5 (20%)	0/2 (CS) (0%)
<b>Kornaat et al<sup>29</sup></b>	CS	–	OR=1.71 (99% CI 0.8 to 3.6)	Age, sex, BMI, intrafamily effects		
Link et al <sup>27</sup>	CS	p>0.05	–	NA		
Fernandez-Madrid et al <sup>22</sup>	CS	NS	–	NA		
<b>Torres et al<sup>19</sup></b>	CS	β=2.50 (95% CI –0.4 to 5.4)	0.82 (–0.5 to 2.1)	Age and BMI		
Bone attrition (level of evidence: conflicting)						
<b>Hernández-Molina et al<sup>27</sup></b>	CS	OR=2.1 (95% CI 1.4 to 3.4)	1.2 (0.7 to 2.0)	Age, sex, BMI, K&L grade, presence of BML and effusion	1/2 (50%)	1/2 (CS) (50%)
<b>Torres et al<sup>19</sup></b>	CS	β=3.3 (95% CI 1.8 to 4.9)	1.9 (0.7 to 3.1)	Age and BMI		

Author's name in bold indicates high-quality studies; 'positive' in front of p values indicates significant positive association sizes. r: (Spearman's or Pearson's) correlation coefficient between MR feature of interest and pain in continuous scale (WOMAC pain subscale or VAS); in a cohort study the correlation coefficient showed the association between changes of the MRI features with the changes in pain during the follow-up. OR, odds of having pain (in cross-sectional studies) or increasing pain (in cohort studies) when a MRI feature is present or increasing comparing to the odds when MRI feature is absent. is regression coefficient representing the increase in knee pain severity associated with increase in lesion score, PR, prevalence (odds) ratio. ACL, anterior cruciate ligament; BMI, body mass index; BML, bone marrow lesion; C, cohort, CS, cross-sectional studies; K&L, Kellgren and Lawrence; LCL, lateral cruciate ligament; MCL, medial cruciate ligament; NA, not applicable; NS, not significant; PCL, posterior cruciate ligament; VAS, Visual analogue scale; WOMAC, Western Ontario and McMaster Scoring.

**Bone attrition**

Conflicting evidence was found on the association between bone attrition and pain. One<sup>19</sup> of two high-quality cross-sectional studies,<sup>19 27</sup> showed a positive association.

**Sensitivity analysis**

When we used median score of all studies instead of mean score as the cut-off of high quality studies, the level of evidence of the association of all MRI finding investigated remained the same. The number of positive studies without quality assessment is shown in table 2.

**DISCUSSION**

Pain is the most disabling symptom of OA. Knowledge about the structures that cause pain is crucial, because in the future it may be possible to specifically target interventions. For a long time, research on the structural cause of pain has been focused on cartilage defects, even though cartilage does not have pain fibres.<sup>3</sup> Further, research on structures that produce pain in the knee was hampered by the limited ability of radiographs to visualise knee structures extensively. MRI has been shown to be superior to plain films. It demonstrates the whole joint organ. Since several initial reports seemed positive about the association between MRI findings and pain, we therefore investigated the evidence

between the MRI findings and knee pain in patients with knee OA. Our findings will be relevant to researchers, clinician and radiologists reporting MRI studies.

We identified a moderate level of evidence for a positive association for BML and effusion/synovitis with pain in knee OA. The level of evidence was limited for a positive association for knee ligamentous abnormalities. We found limited levels of evidence for no association for osteophytes and subchondral cysts. Conflicting levels of evidence were found for cartilage defects, meniscal lesions and bone attrition. We did not investigate studies found during the literature search which investigated features beyond the scope of this review: patella alignment,<sup>37</sup> peripatellar and other periarticular lesions,<sup>38</sup> popliteal or synovial (Baker's cyst).<sup>13 26 29</sup>

In our review, we used a priori defined qualitative levels of evidence to reach a summary. We consider this as a strength because we provide an alternative to quantitative statistics, which could not be calculated as the topic of our review included several aspects of studies that were heterogenic. However, simply counting positive studies also has several drawbacks. It does not take into account the size of the studies, and the decision on 'positive or negative' studies was based only on statistical significance. In meta-analysis, it is theoretically possible that individual studies are negative but the pooled effect is positive.<sup>39</sup> Another technical limitation of our review is the use of quality scores to assess the methodological quality of the studies. It could be that when different quality score sets were used, the interpretation of the results could be influenced.<sup>40</sup> Other limitations of this review mostly reflect the limitations of the studies investigated. First, no publication bias could be assessed using a funnel plot due to the limited number of studies that reported their results in RR or OR.<sup>41</sup> Therefore, we do not know whether preferentially positive findings were published. Second, the quality of included studies was not excellent. There are several obvious examples of limitations of the studies. MRI scan interpretation is by nature subjective, as few, if any, quantitative methods exist. Attempts at standardisation may not be generally used. Also, most scans were read unblinded to order. It is possible that MRI readers define the later findings as more severe than the first findings. This could lead to misclassification.

The moderate associations found in the review have the consequence that more research is needed.<sup>42</sup> Epidemiological studies about BML and effusion/synovitis could strengthen the levels of association. An ideal epidemiological study design would be a case-crossover study where individual MRI findings in the presence of knee pain at one time point are compared with MRI findings in the same patient without knee pain at another time point. The ideal data analysis would give an association size and permit adjustment for confounders, including age and sex, and also for other MRI features when multiple MRI findings are studied simultaneously.

The causal relationship between BML and effusion/synovitis and pain in knee OA needs further study. Our knowledge is now limited to the fact that BML, defined as ill-defined hyperintensities on T2-weighted MRI,<sup>43</sup> comprises normal tissue, oedema, necrosis and fibrosis in histological slices.<sup>44</sup> Further, although knee OA is not considered as an inflammatory arthritis per se, research on the role of inflammation in knee OA and the potential use of anti-inflammatory treatments in knee OA should also be pursued in the light of the possible association between effusion/synovitis with knee pain in knee OA. Evaluation of effusion and synovitis can be improved by using contrast enhancement,

since it can highlight inflammation and improve the distinction between synovitis and effusion.<sup>12 19</sup> Gadolinium contrast diffusion is affected in synovitis tissue, where the blood flow and permeability are changed.<sup>45</sup> In the present review, no included papers performed contrast-enhanced MRI.

Beyond the knee itself further research needs to be focused on the origin of pain in OA and representation in the central nervous system. Some observations have shown that pain in arthritis is also characterised by abnormal pain response (hyperalgaesia)<sup>46</sup> and functional MRI has the potential to study hyperalgaesia and other pain response.

Knowing which structures in the knee are associated with knee OA will add to our understanding of OA and, in the long term, will lead to rational therapeutic targets for OA. This will mean improvement in patient care, since at this moment the therapeutic options against OA are limited.<sup>47</sup> At present, the clinical implication of BML is not clear, despite being a common finding in knee OA, being present in 78% of patients with knee OA with pain and in 30% of patients with knee OA without pain.<sup>24</sup> BML is plainly not pathognomonic of knee OA as it is also found in a range of conditions such as trauma, osteoporosis and rheumatoid arthritis.<sup>48</sup> Moreover, BML is also not a static finding. Almost every BML in knee changes in size over a period of 3 months.<sup>49</sup> The clinical implications of effusion/synovitis may be clearer, since they might permit the potential use of anti-inflammatory drugs in treatment of OA. Effusion/synovitis is common in knee OA. Moderate effusion being seen in 36% of patients with knee OA and synovitis present in (84%) of knees.<sup>26</sup>

The finding that ligamentous abnormalities may associate with pain is of special interest. While the exact aetiology and management of these finding remains unclear it may be that surgical intervention could in theory be aimed at repair of these structures to alleviate pain. However, based on present knowledge, surgical intervention for symptomatic treatment is not currently indicated.

In summary, this systematic review has shown that BML and effusion/synovitis were associated with knee OA pain. However, the level of evidence is moderate and these features need to be explored further.

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**Competing interests** All authors have completed the Unified Competing Interest form at [http://www.icmje.org/coi\\_disclosure.pdf](http://www.icmje.org/coi_disclosure.pdf) (available on request from the corresponding author).

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