Can minimal clinically important differences in patient reported outcome measures be predicted by machine learning in patients with total knee or hip arthroplasty? A systematic review

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Abstract
Objectives: To systematically review studies using machine learning (ML) algorithms to predict whether patients undergoing total knee or total hip arthroplasty achieve an improvement as high or higher than the minimal clinically important differences (MCID) in patient reported outcome measures (PROMs) (classification problem).

Methods: Studies were eligible to be included in the review if they collected PROMs both pre- and postintervention, reported the method of MCID calculation and applied ML. ML was defined as a family of models which automatically learn from data when selecting features, identifying nonlinear relations or interactions. Predictive performance must have been assessed using common metrics. Studies were searched on MEDLINE, PubMed Central, Web of Science Core Collection, Google Scholar and Cochrane Library. Study selection and risk of bias assessment (ROB) was conducted by two independent researchers.

Results: 517 studies were eligible for title and abstract screening. After screening title and abstract, 18 studies qualified for full-text screening. Finally, six studies were included. The most commonly applied ML algorithms were random forest and gradient boosting. Overall, eleven different ML algorithms have been applied in all papers. All studies reported at least fair predictive performance, with two reporting excellent performance. Sample size varied widely across studies, with 587 to 34,110 individuals observed. PROMs also varied widely across studies, with sixteen applied to TKA and six applied to THA. There was no single PROM utilized commonly in all studies. All studies calculated MCIDs for PROMs based on anchor-based or distribution-based methods or referred to literature which did so. Five studies reported variable importance for their models. Two studies were at high risk of bias.

Discussion: No ML model was identified to perform best at the problem stated, nor can any PROM said to be best predictable. Reporting standards must be improved to reduce risk of bias and improve comparability to other studies.

Introduction
Total hip arthroplasty (THA) and total knee arthroplasty (TKA), also referred to as total hip or total knee replacement [1], both subsumed under the term total joint arthroplasty (TJA) [2] reflect a common medical treatment in developed OECD countries. The rates for joint
replacement are increasing rapidly in OECD countries within the last decades. From 2000 to 2013, a 35 percent increase in THAs and a roughly 100 percent increase in TKAs has been reported. However, there exist huge differences in TJA rates across countries within the OECD. Even though age is a strong predictor for the need of TJA, variations in population age are not the key driver of differences in TJA across countries [3]. For OECD countries, the trend in THAs is predicted to grow from 184 implants per 100,000 inhabitants (2015) up to 275 implants per 100,000 inhabitants in 2050 [4]. For TKA, the annual growth rate across OECD countries is estimated to be 5.5 percent per year [5]. Additionally, various international studies predict (highly) increasing rates of TKA and/or THA for countries such as the UK [6], Germany [7, 8], New Zealand [9], Sweden [10, 11], the US [12–14], Australia [15] or Taiwan [16]. However, it has been reported that up to 30 percent of patients undergoing TJA remain unsatisfied [17]. Thus, the question arises whether it is possible to select only patients for surgery that will also be satisfied afterwards.

Over the past decades, various standardized, patient reported measures have been developed to capture patient’s perception of their health or quality of life. These measures are referred to as patient reported outcome measures (PROMs). So far, the implementation of PROMs in clinical practice is rare [18]. There exist both indication/disease/condition specific PROMs such as for TKA [19] or THA [20], but also PROMs that measure generic health status [21]. Once measured before and after a medical treatment has been conducted, PROMs provide the possibility to evaluate whether a clinical treatment has led to an improvement relevant to the patient. Such improvements, specifically the smallest still relevant to the patient, are also referred to as minimal clinically important differences, short MCIDs [22, 23]. MCIDs for PROMs can be derived using different methods such as distribution-based methods [24, 25], anchor-based methods, or by expert consensus [23, 26].

Knowing that a substantial share of patients fail to achieve MCIDs after TJA [2, 27–29], it would be a huge advantage to know which patients will or will not achieve a MCID from a given treatment before the treatment is conducted [2]. If possible, accurate predictions can reduce costs for the healthcare system by preventing patients that will not benefit from treatment from unnecessarily receiving it, decreasing their risk associated with surgery and facilitate optimal resource allocation within the healthcare system.

An approach that could be exploited for predicting whether patients will receive MCIDs after TJA is machine learning (ML), a branch of artificial intelligence (AI) [30]. So far, ML methods have shown to be able to outperform more traditional methods such as regression techniques in various prediction tasks [31–34]. When ML is applied to prediction tasks, typically supervised ML is used [35]. Supervised ML is trained to perform predictions based on various variables (features), using training data where the outcome variables value is known (labelled) [35, 36]. Once trained, supervised ML is used to predict outcomes in unlabeled test data, which contains the same features as the training data. While predicting continuous outcomes in ML is referred to as regression, categorical outcome prediction is said to be classification [35]. However, there are various models available for application in a classification task. Among those, it is not clear a priori which model will perform best on a given task [35, 37]. Instead, researchers rely on trial and error, testing various models regarding their predictive performance, then selecting the model that performs best [35].

In contrast to traditional statistical prediction models such as logistic regressions (LRs), machine learning typically requires less human input, is less theory led and handles nonlinear relationships of variables [37], variable selection or interactions itself [27]. In application, traditional models are rather designed to identify associations than performing predictions [2]. However, there exist various distinctions of ML and traditional models, ranging from classifying anything else than traditional regression as ML [38] to defining the difference between statistical models and machine learning as a continuum, where a model is closer to ML the less human input (e.g. defining interactions, non-linear specifications) it requires [39]. Breiman [40] described ML and traditional models rather as two cultures. One culture aims to predict outcomes with given inputs without the aim to explain the relationship between inputs and outputs in detail. The other culture rather aims to model the relations between input and output correctly but is not deeply interested in achieving best predictive performance. For the reason of this study, we define ML as methods other than traditional LR or linear regression [38, 41], thus included models handle at least non-linear relationships, feature selection or interactions themselves. Consequentially, we distinguish from Christodoulou et al. [42] and also include LASSO (least absolute shrinkage and selection operator) models as ML into our review, setting the cut-off between ML and traditional models closer to the traditional models edge. However, some studies [27, 28] included have also performed LRs. LR results will not be presented in the results section of this paper.

ML techniques have already been applied and systematically reviewed in various prediction tasks in healthcare such as for sepsis prediction [43], psychiatric disorders prediction [41], neurosurgery outcomes [34], therapeutic
outcomes in depression [44], and more. However, no systematic review summarizing the results for ML in the prediction of MCIDs in PROMs for patients undergoing TJA has been conducted so far.

Following the PICOTS scheme [45], our aim was to systematically review studies applying machine learning (I/C) in order to predict, based on pre-surgery data (T) from TKA/THA patients (P/S), whether or not patients that underwent TKA or THA (P) achieve a difference in pre- and post-surgery (T) PROM scores as high or higher as a derived MCID (O) (binary outcome, classification task).

Methods
Protocol and registration
The study was registered in the PROSPERO registry (ID: CRD42021229935) for systematic reviews on 4th of February 2021. No protocol has been published. We followed the Preferred Reporting Items for Systematic reviews and Meta-Analysis of Diagnostic Test Accuracy Studies (PRISMA-DTA) checklist for the structure of this systematic review.

Eligibility criteria
Studies included in this review must satisfy the following criteria: patients included in the study underwent either total knee or total hip arthroplasty (of any etiology), or both; reported at least one PROM both before and at least one time after treatment; written in English or German; MCIDs were derived either anchor-based, distribution-based or through expert consensus (or referenced literature must include either of the three calculation categories); MCID calculation method must have been reported (at least in the referenced literature); predictions for MCIDs after treatment were performed; prediction models were based on machine learning; predictive performance was assessed using either area under the receiver operating curve (AUC)/c-statistic, J-statistic (Youden-Index), G-mean, F1-measure, sensitivity and specificity, or accuracy.

Studies were excluded from this review if they met the following criteria: case studies or reports; books; reviews; congress articles or presentations; only applied traditional statistical models; outcomes were not patient reported.

Information sources
The search was conducted on November 2nd, 2021. The databases MEDLINE, PubMED Central (PMC), Web of Science Core Collection, Google Scholar and the Cochrane Library were searched. The initial search term consisted of several variations of four blocks that build up the search term components (see Additional file 1: Detailed search terms for each database searched). Each block consisted of several variations including MeSH terms, truncations, synonyms, acronyms, or related terms of the component aimed to be identified by each block. The first block consisted of terms related to supervised machine learning, which is the category of method necessary to be applied in the included papers. The second block consisted of terms indicating the use of PROMs, while the third consisted of terms related to MCIDs. Fourth and finally, terms related to total knee or hip arthroplasty were included into the search term.

Search
Search terms are fully available in Additional file 1: Appendix I.

Study selection
After conducting the initial search, all papers identified on all databases were transferred to Citavi 6.7, a literature management software from the Swiss Academic Software GmbH, Wädenswil, Swiss. Next, all duplicates were removed. Further, all articles of excluded document type (see eligibility criteria) were removed. The remaining studies were screened on titles and abstracts. After excluding off-topic papers or papers in the wrong language (see Fig. 1), the rest of studies were read full text. After excluding studies that turned out to be subject to exclusion criteria / did not meet the inclusion criteria, the final papers included into the study were identified. The whole process was conducted by two researchers (BL, AT) independently. Differences in included papers in the different stages (identification, screening, eligibility, and inclusion) were discussed and settled by arguing in line with the inclusion and exclusion criteria. If differences could not be settled, a third researcher (VV) was available for consultation for final settlement. For each stage, we applied a low-threshold strategy. That is, if at least one of the search-conducting reviewers thought it is somehow possible that a paper has hit the inclusion criteria, even if one might assume objectively that it is unlikely, the paper was included for the next stage [46].

Data collection and items collected
Once identified to be included into this systematic review, specific data from studies was collected standardized. That is, a previously constructed, standardized table including features relevant to interpret the predictive performance of the applied models was filled with the data reported by the respective study (Table 1). Specifically, the table includes the following items: Country of data origin, PROMs/MCID values, MCID calculation method, time-difference surgery to post-surgery PROM collection (months), number of observations, number
of features, applied machine learning methods, ratio of training to test dataset, cross-validation applied in the training dataset, whether outlier detection and analysis were performed, whether missing value management was reported, whether feature preprocessing was performed, whether imbalanced data adjustment was performed, the AUC/c-statistic, J-statistic, F1-measure, sensitivity, specificity and accuracy, the Brier score, the best predictive model and best predicted PROM, and the predictive task (classification or regression).

**Risk of bias assessment**
Risk of bias assessment was conducted using PROBAST (Prediction model Risk of Bias ASsessment Tool). PROBAST includes a four-step approach. While steps one and two consider the review question and
Table 1  Extracted data of the included studies

| Study | Fontana et al. [2] | Harris et al. [27]* | Huber et al. [28]* | Katakam et al. [59] | Zhang et al. [60] | Kunze et al. [29] |
|-------|-------------------|---------------------|-------------------|---------------------|------------------|------------------|
| Country of data origin | US | US | UK | US | Not reported | US |
| Surgical procedure | THA/TKA | TKA | THA/TKA | TKA | TKA | THA |
| PROMs/MCID values | HOOS JR: 17.7 KOOS JR: 13.6 SF-36 (MCS+PCS): 5.0 (both) | KOOS Total: 91.8 KOOS JR: 20.8 KOOS Pain: 25.0 KOOS Symptoms: 14.3 KOOS ADL: 24.6 KOOS Quality of Life: 12.5 KOOS Recreation: 17.5 | EQ VAS Hip: 11 EQ VAS Knee: 10 OHS: 8\textsuperscript{a} OKS: 7\textsuperscript{a} | KOOS: MCID value not reported PROMIS Global PF: MCID value not reported | PROMIS Global MH: MCID value not reported | EQ VAS: Not reported |
| MCID calculation method | Anchor-based | Distribution-based | Anchor-based | Distribution-based (VAS) Anchor-based (OKS, OHS) | Distribution-based | Anchor-based Distribution-based\textsuperscript{b} |
| Time-difference surgery to post-surgery PROM collection (months) | 24 | 12 | 12 | 12 | 24 | 24 |
| Number of observations | 7,239 (THA) 6,480 (TKA) | 587 | 30,524 (THA) 34,110 (TKA) | 744 | 2840 | 616 |
| Number of features\textsuperscript{c} | 66–97 | 6–106 | 81 (candidate predictors) | 24 (candidate predictors) | 18 (WOMAC); 19 (other PROMs) | 8 |
| Applied machine learning methods | LASSO Random forest Support vector machine | LASSO Gradient boosting machine Quadratic discriminant analysis | Extreme gradient boosting machine Random forest Multistep adaptive elastic net Neural network Naive Bayes k-nearest neighbours Boosted logistic regression | Stochastic gradient boosting Random forest Support vector machine Neural network Elastic-net penalized logistic regression | Support vector machine LASSO Random forest Extreme gradient boosting | Stochastic gradient boosting Random forest Support vector machine Neural network Elastic net penalized logistic regression |
| Ratio of training to test dataset | 80.20 | No test dataset | About 1:1 (dataset of the next year) | 80.20 | 80.20 | 80.20 |
| Cross-validation applied in the training dataset | Yes | Yes | Yes | Yes | Yes | Yes |
| Outlier detection and analysis performed? | Not reported | Not reported | Not reported | Not reported | Not reported | Not reported |
| Missing value management reported? | Yes | Not reported | Yes | Yes | Yes | Yes |
| Feature preprocessing performed? | Yes | Not reported | Not reported | Not reported | Not reported | Not reported |
| Imbalanced data adjustment performed? | Not reported | Not reported | Yes | Not reported | Yes | Not reported |
| AUC/c-statistic\textsuperscript{d} | 0.89 (not reported) | 0.72 (not reported)\textsuperscript{d} | Not reported on test data | 0.77 (0.74–0.79) | SVM: 0.95 (0.94–0.97) XGB: 0.95 (0.94–0.97) | 0.97 (0.94–0.99) |
| J-statistic | – | – | 0.59 (not reported) | – | SVM: 0.85 (not reported) XGB: 0.86 (not reported) | – |
| F1-measure | – | – | 0.78 (not reported) | – | – | – |
classification of the prediction model, step three assesses the risk of bias (ROB) and applicability, and finally step four closes with an overall judgement [47]. The assessment tool assesses participants, predictors, outcomes and analysis ROB as well as participants, predictors and applicability to the review question [45]. For PROBAST assessment, risk of bias is thus assessed both at study and outcome level.

### Summary measures
Outcomes of studies needed to be reported as AUC/c-statistics, J-statistics, G-mean, F1-measure, sensitivity and specificity, or accuracy.

### Synthesis of results
Due to the variety of models, features and included PROMs from which outcomes (MCIDs) were derived in the studies, no meta-analysis could be conducted.

### Results

#### Study selection
Searches of the search terms were conducted on the previously mentioned databases (see section “Data sources”) on November the 2nd, 2021. Searches in PubMed Central, MEDLINE, Web of Science Core Collection, Google Scholar and Cochrane Library yielded 139, 355, 314, 237 and 100 hits, respectively, resulting in 1145 hits in total.

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**Table 1** (continued)

| Study          | Fontana et al. [2] | Harris et al. [27]* | Huber et al. [28]* | Katakam et al. [59] | Zhang et al. [60] | Kunze et al. [29] |
|----------------|--------------------|--------------------|--------------------|--------------------|-------------------|-------------------|
| Sensitivity    | –                  | –                  | 0.82 (not reported) | –                  | SVM: 93.1 (not reported) | XGB: 95.6 (not reported) |
| Specificity    | –                  | –                  | 0.77 (not reported) | –                  | SVM: 86.8 (not reported) | XGB: 84.9 (not reported) |
| Accuracy       | –                  | –                  | 0.79 (not reported), balanced accuracy | –                  | –                  | –                  |
| Brier Score\(^b\) | Not reported       | LASSO (KOOS Pain): 0.16 (not reported) | LASSO (KOOS Symptoms): 0.17 (not reported) | XGB: 0.16 (not reported) | SVM: 0.12 (not reported) | XGB: 0.11 (not reported) |
| Best predictive model | Logistic LASSO | Random forest | LASSO, Gradient boosting machine, QDA (Pain) | Extreme gradient boosting | Neural Network Elastic-net penalized logistic regression | Support vector machine (SVM) | Random forest |
| Best predictive PROM | SF-36 MCS | KOOS Pain | KOOS Symptoms + KOOS ADL | EQ VAS (hip) | KOOS | SF-36 MCS | EQ VAS |

| Predictive task | Classification | Classification | Classification | Classification | Classification |

HOOS JR, Hip disability and osteoarthritis outcome score joint replacement; KOOS JR, Knee injury and osteoarthritis outcome score joint replacement; SF-36 MCS, Short form-36 mental component score; SF-36 PCS, Short form-36 physical component score; EQ, EuroQol; VAS, Visual analog scale; OKS, Oxford Knee Score; OHS, Oxford Hip Score; LASSO, Least absolute shrinkage and selection operator; AUC, area under the receiver operating curve; QDA, Quadratic discriminant analysis; ADL, Activities of daily life; JR, Joint replacement

\(^a\) Also applied LR
\(^b\) Value was taken from literature
\(^c\) This value was calculated on postoperative score distribution
\(^d\) Finally included in the models when not otherwise stated
\(^e\) Result from the training dataset with fivefold cross validation as no AUC was reported on test data
\(^f\) Confidence intervals (95% if not otherwise specified) in parenthesis
All hits were transferred to Citavi 6.7. 610 unique titles were identified. 93 were excluded based on document type (see eligibility criteria). After screening title and abstract of the remaining 517 studies, 18 studies underwent full text screening. Thereof, six titles were included after full-text reading (see Fig. 1).

From the 18 articles screened for full text, two studies were excluded due to including the wrong study population from the perspective of this review. One study [48] included knee arthroscopies and one [49] patients with osteochondral allograft for cartilage defects. Next, five studies [50–54] did not calculate MCIDs, even though they researched PROMs in TJA patients. Finally, six studies [26, 55–58] aimed to make comparable predictions as searched for this review except failing to apply machine learning.

Study characteristics

Machine learning models applied
The studies included applied various machine learning techniques, while two of them [2, 27] also included LRs. One for comparison with ML [27], the other defined LR as ML [28]. The number of included ML techniques varied from three [2] to seven [28]. Overall, the four most commonly applied algorithms across studies were random forest [2, 28, 29, 59, 60], gradient boosting machine (GBM) \(^1\) [21, 28, 29, 59, 60], support vector machine (SVM) [2, 29, 59, 60] and LASSO [2, 27, 60]. We reported, for each study, the model that performed best on the test dataset on the studies main outcome metric, which was the AUC for all studies except for Huber et al. [28], where J-statistic was the favored outcome on the test dataset. In case only a validation dataset was used and no test dataset [27], the main outcome on the validation dataset was reported. For the model(s) performing best on the main outcome, we also reported all other outcomes reported in the studies.

Sample size, data origin and number of features
Sample size varied widely across studies. While Harris et al. [27] only included 587 TKAs (both for training and validation) in their analysis, Huber et al. [28] included 34,110 individuals with TKA for model development and 34,406 TKAs for testing. For THA, they used 30,524 and 31,905 individuals for training and testing, respectively. The other studies ranged in between (see Table 1). While Huber et al. [28] used data from NHS treated patients among patients treated in multiple centers (all run by NHS), Katakam et al. [59] used data from five sites, Harris et al. [27] included three sites (VA medical centers), Zhang et al. [60], Kunze et al. [29] and Fontana et al. [2] only exploited data from one site.

Due to the different datasets utilized, the number of features initially available to the models varied across studies. Fontana et al. [2] had access to 51 initial variables in their “before surgery” setting. However, as 25 of them were categorical, the number of features (including the categories dummies) included in their analysis must have been higher but was not reported. Further, the number of features used by each model was not reported. Harris et al. [27] came up with 106 variables out of which models were able to select the variables relevant for predictions. Their final models included six to 106 variables, depending on algorithm and PROM. Huber et al. [28] had access to 81 variables out of which models were able to select their individual number of predictors, however the final number of features per model was not reported. Kunze et al. [29] initially had access to eleven variables to be included in predictive models. After recursive feature selection, they came up with eight variables included into the final models. Katakam et al. [59] initially had access to 24 variables (including dummies for categorial variables). They did not report the final number of variables after feature selection exploiting the random forest. Finally, Zhang et al. [60] included 18 variables for the WOMAC and 19 variables for the other PROMs in their final prediction models.

Training, validation, and testing
All included studies used some approach to perform training and validation of their developed models. However, studies applied different approaches to do so. Harris et al. [27] performed cross-validation with bootstrapping on the training dataset to assess their models’ performance (internal validation) but did not exploit any unforeseen test dataset. Huber et al. [28] used cross-validation in the training dataset for model selection and tested their models on an independent test dataset of the subsequent year. The remaining studies split their dataset beforehand randomly into training and test data. While the proportions of training to dataset were 20:80 in Fontana et al. [2], Kunze et al. [29] and Zhang et al. [60], Katakam et al. [59] split the dataset into 70 percent training and 30 percent test data. During model training, all studies applied cross-validation to perform hyperparameter tuning and model selection. Then, performance evaluation was executed on the test dataset which was completely unforeseen by the algorithms. Huber et al. [28] and Zhang et al. [60] applied upsampling (a replication of the minority class to receive a balanced training dataset) in the training dataset to account for disproportions across outcome groups.

\(^1\) Note that several GBM variations such as stochastic gradient boosting, extreme gradient boosting or gradient boosting are summarized under the umbrella of GBM.
**Predictive performance**

To evaluate models’ predictive performance, papers reported different performance indicators. For performance comparison in this section, only performance reported on the test dataset is included since it is indicative of model generalization, while performance assessment on training data could be biased due to overfitting. That is, models fit the training data disproportionately well but have poor generalization on test data [35]. As an exception, in case of development studies, performance on training data was reported in case cross-validation was applied to account for overfitting.

Most of the studies reported the AUC as main performance measure. The AUC is calculated as the area under the receiver operator curve, and the receiver operator curve is a plot of sensitivity against false positive rate (1-specificity) of a given predictive tool using different decision thresholds for categorizing outcomes as either positive or negative [61]. AUC/c-statistic values are classified as fail (0.5–0.59), poor (0.6–0.69), fair (0.7–0.79), good (0.8–0.89) or excellent (0.9–1.0) [27, 62]. The best model in Harris et al. [27] and Katakam et al. [59] performed fair, the best model in Fontana et al. [2] good and the models of Kunze et al. [29] and Zhang et al. [60] excellent on the test/validation sample (see Table 1). Unfortunately, Huber et al. [28] did not report AUC results for the test sample. Applying their models to the test sample, they came up with a J-statistics of 0.59. The J-statistics (Youden Index) is the sum of sensitivity and specificity minus one [63]. If the proportion of predicted positives in the true positives group is higher than in the true negatives group, the value is always above zero [63].

**Utilized PROMs and MCID derivation**

PROMs used varied widely across studies. Across all included studies, sixteen different PROMs have been applied to KA and six to HA patients, respectively. No PROM was utilized in all studies. All included studies except Kunze et al. [29] included at least three PROMs. Kunze et al. [29] only included a generic PROM. Harris et al. [27] only included treatment-specific PROMs, the remaining studies included both generic and treatment specific PROMs.

Fontana et al. [2] included four PROMs in their paper, namely SF-36 physical component score (PCS), SF-36 mental component score (MCS), Hip Disability and Osteoarthritis Outcome Score for Joint replacement (HOOS JR) and the Knee Disability and Osteoarthritis Outcome Score for Joint replacement (KOOS JR). While the former are generic health status scores with either an additional focus on physical (PCS) or on mental health (MCS) [64], the latter are knee or hip specific scores [65]. Harris et al. [27] included KOOS Total, JR and the subscales KOOS pain, symptoms, activities of daily living (ADL), quality of life (QoL) and recreation. Huber et al. [28] included the EQ-5D-3L and EQ VAS (both general health) as well as the Oxford Knee Score (OKS) and Oxford Hip Score (OHS), which are both disease specific, whereas Kunze et al. [29] only included the generic EQ VAS. Katakam et al. [59] included the KOOS as disease-specific PROM and the Patient Reported Outcomes Measurement Information System (PROMIS) Global PF, PROMIS Global MH [66] and numerical rating scale for pain (NRS Pain). Finally, Zhang et al. [60] reported the Western Ontario and McMaster Universities Osteoarthritis Index (WOMAC) as well as SF-36 PS and SF-36 MCS.

MCIDs can be calculated either anchor-based, distribution-based or based on evidence from previous studies [67]. All studies used distribution-based or anchor-based methods or referred to such. While for HOOS JR and KOOS JR they adopted values based on anchor-based methods from the literature, Fontana et al. [2] determined the score for SF-36 PCS and MCS themselves based on distribution-based methods. Harris et al. [27] calculated MCIDs applying anchor-based methods, with the Self-Administered Patient Satisfaction Scale (SAPS) as anchor. To determine the score for each PROM which discriminates patient satisfaction best, they used the Youden index. Huber et al. [28] calculated MCIDs using distribution-based methods (half a standard deviation) for EQ VAS (preoperative score) and referred to the literature for OHS and OKS, respectively. Kunze et al. [29] also defined MCIDs based on distribution-based methods as half of a standard deviation. However, in contrast to the other studies, they used postoperative scores instead of preoperative scores to determine MCIDs. Katakam et al. [59] derived all MCIDs by distribution-based methods. Zhang et al. [60] referred to the literature which based the MCID calculation on anchor-based methods for MCID derivation of all PROMs.

**Variable importance**

Out of the six included studies, five [2, 28, 29, 59, 60] reported variable importance. If studies reported variable importance for different algorithms, for simplicity, only the variable importance for best performing algorithms as well as the best predictive PROM will be reported. Among those, preoperative PROM scores were the most predictive variables in all studies. In four studies [2, 28, 59, 60], a depression-indicating variable was among the top five most important predictors. Further, three studies [2, 28, 59] had at least two other PROMs or PROM subscales ranging among the top five predictors for MCIDs. As Kunze et al. [29] did not include more than one PROM in their study, it was not possible to have other PROM scores among the best predictor variables.
Missing data

Studies facilitated different strategies to deal with missing data. Fontana et al. [2] handled numeric missing variables by imputation to the mean, while for categorical variables, an extra class was created for missing values, exploiting information of missing values. Huber et al. [28] removed all patients with missing values and variables with variance close to or at zero. Kunze et al. [29] and Katakam et al. [59] performed multiple imputation for variables with less than 30 percent missing values. Kunze et al. [29] excluded one variable with more than 30 percent missing values. Zhang et al. [60] reported two variables with few missing values and applied imputation with mean values. Harris et al. [27] did not report missing values. A detailed overview of missing values for each study is given in Additional file 2: Missing data values for included studies.

Risk of bias assessment

Risk of bias assessment within studies was conducted using PROBAST (Prediction model Risk of Bias ASsessment Tool), which assesses the bias for the prediction tools. PROBAST includes a four-step approach. While steps one and two consider the review question and classification of the prediction model, step three assesses the risk of bias (ROB) and applicability, and finally step four closes with an overall judgement [47]. Table 2 represents the suggested tabular presentation for PROBAST results by Wolff et al. [47]. A guidance on how to perform PROBAST is also given at the website probast.org, and in Moons et al. [45] (Additional file 3: Probst assessment of all included studies).

After conducting the PROBAST assessment, four studies were free of ROB, while two were of high risk of ROB. All studies were applicable, that is, populations, predictors and outcomes fit this reviews purpose [47]. All studies had in common that predictors were not excluded from the outcome definition, yielding a possible bias introduction [45]. As the case in this setting, the outcome (MCID) is partially defined by pre-surgery PROMs, a predictor also applied in the prediction models. Therefore, bias was not assumed to be introduced by pre-operative PROMs due to the nature of the study setting. Next, all studies determined the outcome with knowledge of predictor information, i.e. predictors were known as the outcome was determined [45]. That is inherent to the study setting and is not assumed to introduce any bias. Further, almost all studies had events per variable (EVP) – that is, the number of participants with the least often reported outcome (MCID vs. no MCID) over the number of candidate predictors – below the recommended value for machine learning of 200 by Moons et al. [45]. However, Moons et al. [45] refer to van der Ploeg et al. [68] for the value of 200. van der Ploeg et al. [68] indicated a number of EVP ≥ 200 if studies AUC was reported on the training dataset and not tested on a validation dataset, calculating the bias (or optimism) as difference between the AUC on the training and on the validation dataset. As stated, all studies at least applied cross-validation to account for overfitting, thus no study reported outcome metrics only on the training dataset without any validation [68]. Consequently, EVP was not considered to be able to introduce bias in all studies.

However, two studies [27, 28] were subject to risk of bias beyond that common traits. Harris et al. [27] did not report how missing values were handled, nor if there were any, resulting in a high ROB judgement. Note that, following Moons et al. [45], Harris et al. [27] can only be rated as model development study but not as validation study, as they only performed internal validation (cross-validation), but no validation on an external test dataset [45].

Huber et al. [28] did not report calibration of models as outcome metric, a potential ROB in the analysis domain [45]. Additionally, important metrics such as AUC, which

| Study          | ROB Participants | ROB Predictors | ROB Outcome | ROB Analysis |
|---------------|------------------|---------------|-------------|--------------|
| Fontana et al. [2] | +                | +             | +           | +            |
| Harris et al. [27] | +                | +             | −           | +            |
| Huber et al. [28] | +                | +             | −           | +            |
| Katakam et al. [59] | +                | +             | +           | +            |
| Zhang et al. [60] | +                | +             | +           | +            |
| Kunze et al. [29] | +                | −             | +           | +            |

Table 2: PROBAST ROB and applicability assessment results* for all included studies following the suggested tabular presentation by Wolff et al. [47].

ROB, Risk of bias

* + indicates low ROB/low concern regarding applicability; − indicates high ROB/high concern regarding applicability; and ? indicates unclear ROB/unclear concern regarding applicability.

The table above shows the risk of bias (ROB) and applicability assessment results* for all included studies following the suggested tabular presentation by Wolff et al. [47]. The table is structured to reflect the PROBAST assessment criteria, with columns indicating ROB for participants, predictors, outcome, and analysis, as well as applicability for participants, predictors, outcome, and overall ROB and applicability. The study by Fontana et al. [2] is assessed as having low ROB and low concern regarding applicability, while Harris et al. [27] and Huber et al. [28] are rated as having high ROB and high concern regarding applicability. The study by Kunze et al. [29] is assessed as having unclear ROB and unclear concern regarding applicability.

As shown in the table, all studies were applicable, meaning that populations, predictors, and outcomes fit the review’s purpose. This indicates that the studies are suitable for the review’s objectives.

The absence of bias in the studies is crucial for the validity of the review’s findings. The presence of ROB, as indicated by high ROB, signifies that the studies may have introduced bias, potentially affecting the reliability of the results. The unclear ROB classification indicates uncertainty in the bias assessment, necessitating caution in interpreting the findings.

The inclusion of missing data handling strategies is also highlighted, reflecting the importance of addressing data missingness in studies. Strategies such as imputation or exclusion of missing values were employed, ensuring a consistent approach across the studies.

In conclusion, the PROBAST assessment provides a comprehensive evaluation of the included studies, ensuring that the review’s findings are based on studies that meet the necessary criteria for validity and applicability. This underscores the importance of rigorous methodology in systematic reviews to ensure the reliability and generalizability of the results.
is commonly reported across all other studies included in this review, was only reported on the training dataset. However, on the test dataset, it was not reported, and it was further not stated why it was not reported on the test dataset. Furthermore, Huber et al. [28] dropped all participants with missing values, another potential for bias introduction [45]. Moreover, Huber et al. [28] did only imprecisely describe the study population. Instead of TKA/THA patients, which are the patients the NHS England PROMs dataset consists of [69], this study only reported their participants to be knee or hip replacement patients. The reader unfamiliar with the dataset may conclude that the dataset includes hip/knee replacement patients other than TKA/THA. Consequentially, Huber et al. [28] was characterized as being at high ROB.

Discussion

This paper was the first to systematically review approaches predicting MCIDs in patient reported outcome measures for patients undergoing total hip or knee arthroplasty. As summarized, all six included papers were published within the last two years. Given that a substantial amount of patients undergoing joint replacement remain unsatisfied afterwards [17] and/or do not achieve an MCID in general reported health [28, 29] or condition-specific PROMs [27], there is a need of creating, evaluating, and implementing approaches to accurately identify patients which would remain unsatisfied after surgery. The actuality of papers published indicates that the problem is recently getting attention and that modern approaches like ML are being exploited to identify such patients.

All studies included various models, and models were partially common among studies (mainly random forest, GBM, LASSO, SVM, neural networks, elastic net LR). However, no type of model clearly outperformed others across studies. This is in line with Hastie et al. [35], who state that there is no ML algorithm that is known a priori to perform best on a given problem and thus only application will show which algorithm (with which tuning parameters) is superior in the specific setting. For example, random forest was applied in five studies, but only performed as best model in two. However, Kunze et al. [29] and Katakam et al. [59] described that they performed feature selection with a random forest. It could be that predictive performance was biased in these studies so that the random forest ended up with features best fitting for that method, but other models could have performed better with other features. Nevertheless, among all studies, performance of the applied ML models did not differ too much. Interestingly and counterintuitively, neither the number of features included, nor the sample size seemed to influence predictive performance to a large extent. Specifically, the best predictive model on AUC [29] was developed with the second smallest sample (n = 616) and with the fewest number of features available compared to all other studies. Overall, ML models’ performance was well in the respective prediction task, indicated by the fact that three out of five studies that reported AUC on test data reached good or excellent performance [27, 29, 60]. The AUC is an appropriate metric for measuring discrimination and to our knowledge the only metric applicable to imbalanced data without being biased [70] – a highly valuable characteristic in datasets with MCIDs as outcomes. Besides that, it has its disadvantages. Specifically, in practice, it is often necessary to have outcome metrics that reflect the performance of an algorithm at a specific sensitivity specificity trade-off [71]. In that context, other metrics are more appropriate and might be reported on a balanced test dataset in case of imbalanced data [70]. Four out of six studies [2, 27, 29, 59] did only report the AUC as discrimination metric, while one [28] did completely rely on other metrics on test data. This reflects potential for improvement. The AUC is an important performance indicator and therefore should be reported by all studies. Common metrics such as F1-measure, G-mean or J-statistic could be additionally reported without much extra effort, and we encourage authors to do so.

PROMs included differed widely across studies. Due to the heterogeneity of datasets and PROMs, conclusions about which PROM can be predicted best are not possible yet. However, EQ VAS [28, 29] and SF-36 MCS [2, 60] were each two times the best predictable PROMs in the six included studies. Interestingly, Fontana et al. [2] and Zhang et al. [60] found that MCIDs for SF-36 MCS, a mental health PROM, could be better predicted than somatic ones (HOOS JR, KOOS JR, WOMAC, SF-36 PCS) in TJA patients. In general, there is a certain advantage of using generic PROMs like EQ VAS or EQ-5Ds over indication-specific PROMs such as KOOS or HOOS. With generic (i.e. disease unspecific) PROMs, medical outcomes of procedures such as surgeries for TKA or THA can be made comparable with other healthcare interventions, facilitating cost-effectiveness analysis and improving decision makers information on allocating healthcare resources [72].

MCID calculations for PROMs were performed according to established methods. However, two studies [29, 59] did not report the MCIDs threshold value. This should be a standardized procedure. Reporting MCID values is relevant because then they can be compared with MCID values from other studies. Recent evidence suggests that MCIDs for PROMs are highly variable and can differ substantially across study populations, calculation methods or even within the same calculation method (e.g.
anchor-based calculation, but using different anchors might yield different results) [73]. It remains unclear whether the MCID calculation method has an impact on ML performance to predict MCIDs in total joint arthroplasty patients. This is a relevant question and should be subject to further research. That being said, three studies [2, 28, 29] defined at least some MCIDs based on distributions. Even though distribution-based methods are common practice, it is not clear that they detect differences that really matter to patients rather than being an arbitrary set threshold based on observed data. MCIDs calculated with anchor-based techniques may be preferred as they stronger focus on the patients perspective [74].

Additionally, studies should report all relevant outcome measures, including calibration. For one study [28], the lack of reporting calibration was one factor for categorizing it as high ROB. Further, even if not a source of bias in the ROB [45], confidence interval reporting is highly important as it helps to understand the certainty of the estimates. However, we found that only half of the studies reported confidence intervals at least for the main outcome, showing high potential for improvement. Furthermore, only two studies [28, 60] performed imbalanced data adjustment. This might be crucial when using other performance metrics than the AUC [70] or when models are sensitive to balanced datasets during training [28]. We therefore recommend to at least test balancing classes and report it, even if it yields no benefit. The same holds for feature preprocessing (e.g. variable transformations), a description of the country where the study took place (not reported in Zhang et al. [60]), a precise description of the study population (imprecise in Huber et al. [28]) and reporting how missing data was handled. No study reported how outliers were handled. We advise all machine learning algorithm developers to assess their studies using the IJMEDI checklist for machine learning applications in medicine as described in Cabitza et al. [75]. It helps to avoid such problems. All studies but one [27] properly applied model validation, feature selection and hyperparameter tuning on the training dataset and reported their final results on an unforeseen test dataset, even though one study [28] did not report all important metrics on the test dataset. However, the result that 5 of 6 studies applied their models to independent test datasets provides strong evidence that the predictive performance in forecasting whether patients will receive an MCID or not after TJA is reasonable to believe. Additionally, all papers discussed the clinical utility and limitations of their applications. Two [27, 28] compared their ML models with traditional approaches, Zhang et al. [60] discussed the lack of comparison with traditional models as a limitation.

Two studies [27, 28] had a high risk of bias. Main problems in the studies with high risk of bias could have been easily handled through transparent and reasonable reporting [35]. Further, given that ML prediction models performance is typically assessed using AUC [41, 43, 76], studies should apply this metric to increase transparency and comparability with other studies’ results. Especially when used as the main metric for assessing performance in the training dataset [28], it should not be omitted in the test dataset.

However, this study comes with limitations. First, as a result of the limited number of studies included for the purpose of this review, and due to the results of ROB both within and across studies, the performance of ML models in the stated context based on this body of evidence should be interpreted with caution. Second, we did not search the literature database Scopus due to a lack of access and might have potentially missed relevant studies. Fortunately, evidence suggests high precision of our used search engines in identifying relevant literature [77, 78]. Third, as model parameter tuning is up to the researcher [35], it can always lead to inferior performing models. We were not able to control hyperparameter tuning in the studies and therefore rely on the subjective performance of the individual researchers of the studies in doing so. This also holds that different methods might outperform others when tuned more sophisticated. Therefore, we also recommend reporting various specifications of the used models with the best tuned model indicated. Fourth and finally, we did not compare machine learning methods to other prediction methods. It might be that other prediction tools such as logistic regression or pre-surgery PROMs themselves might perform as good, better, or worse than machine learning. This needs further investigation. ML studies may therefore include other methods as baseline/comparison models.

Conclusion

Given the promising results of models’ performance of the included studies, ML-based applications to support informed decision making as well as to implement an objective instance in shared decision making between clinicians and patients undergoing TKA or THA should be considered for practical implementation. Discussed issues of risk of bias and underreporting must be eliminated in future research to derive transparent and unbiased results. Especially, important metrics such as the AUC, calibration and uncertainty should be reported standardized across and consistently within studies for better comparability. Further, dataset preparation especially with respect to an unforeseen dataset is crucial when it comes to performance assessment and should be
applied in every study. Relevant checklists are available to ensure high quality of studies applying ML in healthcare.

Supplementary Information

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Additional file 1. Appendix 1: Search Terms.
Additional file 2. Appendix 2: Missing data per variable for all included studies.
Additional file 3. Appendix 3: PROBAST assessment of all included studies.

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None.

Authors’ contributions

BL came up with the idea and discussed the structure and the aim of the paper with W. BL and AT both conducted the literature review and study selection as well as the risk of bias assessment independently. VV was available for discussions of study selection as well as the risk of bias assessment independently. W-V was available for discussion and commented by both AT and VV. BL then adopted the comments from AT and W-V and finalized the final draft. All authors read and approved the final manuscript.

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