Machine learning in predicting extracapsular extension (ECE) of prostate cancer with MRI: a protocol for a systematic literature review

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ABSTRACT

Introduction In patients with prostate cancer (PCa), the detection of extracapsular extension (ECE) and seminal vesicle invasion is not only important for selecting the appropriate therapy but also for preoperative planning and patient prognosis. It is of paramount importance to stage PCa correctly before surgery, in order to achieve better surgical and outcome results. Over the last years, MRI has been incorporated in the classical prostate staging nomograms with clinical improvement accuracy in detecting ECE, but with variability between studies and radiologist’s experience.

Methods and analysis The research question, based on patient, index test, comparator, outcome and study design criteria, was the following: what is the diagnostic performance of artificial intelligence algorithms for predicting ECE in PCa patients, when compared with that of histopathological results after radical prostatectomy.

To answer this question, we will use databases (EMBASE, PUBMED, Web of Science and CENTRAL) to search for the different studies published in the literature and we use the QUADA tool to evaluate the quality of the research selection.

Ethics and dissemination This systematic review does not require ethical approval. The results will be disseminated through publication in a peer-review journal, as a chapter of a doctoral thesis and through presentations at national and international conferences.

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INTRODUCTION

Prostate cancer (PCa) is one of the most frequent cancers in men, with about 6600 new cases diagnosed per year.1 Despite its increasing incidence, a global mortality decline has been observed, in most world regions.2 The detection of low-stage cancers (≤T2), with a better prognosis, related to the increasing incidence of the tumour and proactive screening, may contribute towards a decline in the overall mortality trend, along with improvements in PCa treatment, which is unrelated to the number of cases diagnosed.2

Low-stage cancer includes PCa confined to the prostate gland. Local advanced disease includes extraprostatic extension, which is defined as cancer located outside the confines of the prostate, often mixed with adipose tissue. It may also include seminal vesicle involvement. It is very useful to differentiate T2 (organ confined PCa) from T3 (local advanced disease) in order to differentiate patients who may benefit from radical surgery.

Radical prostatectomy has been the gold standard for surgical treatment for organ confined PCa.3 Robotic-assisted radical prostatectomy introduced a minimally invasive surgical treatment for PCa and has been gaining acceptance for surgeons and patients.4

Strengths and limitations of this study

⇒ To our knowledge, this is the first systematic review that aims to comprehensively synthesise the available evidence about the diagnostic performance of artificial intelligence (AI) and machine learning algorithms in predicting extra-capsular extension (ECE) in patients with prostate cancer.

⇒ This protocol has been developed following the guidance of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses Protocols and has been registered with the International Prospective Register of Systematic Reviews.

⇒ We will include all studies with MRI criteria and AI models to detect ECE on pathology, excluding cross-sectional studies, case series, case reports, case-control studies, systematic reviews, conference proceedings and masters or PhD thesis.

⇒ The novelty of AI in the field of radiology, specially in the field of MRI for the diagnosis and staging of prostate cancer, and also the lack of robust studies with great number of cases, may hamper meaningful conclusions for clinicians.
The detection of extraprosthetic extension prior to treatment, in patients with PCa, affects the cancer prognosis. Positive surgical margin is defined as cancer cells touching an inked surgical margin on the pathological specimen. Extracapsular extension (ECE) is a cause of high frequency of PMS, biochemical recurrence, metastatic disease and lower cancer-specific survival after radical prostatectomy.4–8

So far, to predict the risk of advanced disease, clinicians often use staging nomograms. These nomograms, as D’AMICO or CAPRA, are based on prostate specific antigen results in blood test, biopsy Gleason score and clinical T stage on digital rectal examination; additionally, patient’s age and percentage of positive biopsy cores are also included in European patients for CAPRA classification.9–10

Several studies demonstrated an improved accuracy in predicting ECE (pECE), by incorporating information from MRI. The combined use of MRI and clinical risk further improves prediction of pathologic ECE.11

MR scanners able of acquiring high spatial resolution imaging, along with functional sequences (perfusion, DWI), paved the way to the so-called multiparametric MRI (mpMRI) that has radically changed the non-invasive perception for primary diagnosis and staging of prostate carcinoma.12 MpMRI is a well-established imaging modality in PCa assessment, particularly for depicting clinically significant PCa and improves the yields of transrectal US-guided biopsy.13–16 MpMRI is also an accurate method for local staging of PCa, as proven by a recent meta-analysis, but with a heterogeneous sensitivity across the several studies.17

Current mpMRI criteria, used to detect and pECE on pathology analysis, pathologic lymph nodes status and local or biochemical recurrence (BCR), are mostly qualitative with high reader variability, owing to strong interpreter dependency and experience.17 These include capsular contact length, capsular irregularity and/or bulging, obliteration of rectoprostatic angle, asymmetry of the neurovascular bundles, invasion of the periprostatic fat and seminal vesicle invasion.18 Some studies that applied confidence features have not been described.19

And validated algorithms to pECE, obtained from mpMRI will be developed.28 As far as we know, clinically accepted and validated algorithms to pECE, obtained from mpMRI features have not been described.

OBJECTIVES
The aim of this systematic review is to analyse the AI and Radiomics models for detecting ECE and helping the surgeon to be potentially more proficient in terms of surgical outcome. Concurrently, it is also important to analyse if the models using advanced tumour segmentation and ML capabilities could improve accuracy to detect ECE in patients that underwent prostatectomy.

METHODS AND ANALYSIS
The protocol and completed review will be reported following the guidance of the Preferred Reporting Items for Systematic Reviews and Meta-ana...
for Systematic Reviews and Meta-Analyses (PRISMA) Protocols and the PRISMA, respectively.

Eligibility criteria
Studies will be included in this review if they include:
- Adult men patients (>18 years of age), with PCa with a Gleason score>6 on presurgical prostate biopsy, and submitted to MRI before they were operated by radical prostatectomy with pathologic examination.
- MRI Human studies done in MRI scanner with 1.5T or 3T (Tesla) equipment.
- Studies with patients who underwent hormonal or radiotherapy treatment before surgery or with patients undergoing hormonal, focal therapy or radiotherapy treatment to treat PCa, are excluded.
- Human studies done on scanners with less than 1.5T equipment are also excluded. We include all studies using AI algorithms, including ML algorithms based on MRI images to predict ECE on pathology.

The main outcome of this systematic review is the pathology detected ECE in patients with PCa who underwent radical prostatectomy. The objective is the detection of MRI imaging variables that could predict ECE, before surgery.

Study design
Eligible studies will be prospective cohort studies or randomised controlled trials (RCTs) with prognostic factor analysis, published in peer-reviewed journals. All the studies will only be included if information regarding PCa MRI staging and pathological PCa staging is available in the published report. The studies will have to identify MRI criteria and AI models to detect index lesion, ECE and patient’s outcome who underwent prostatectomy.

Cross-sectional studies, case series, case reports, case-control studies, systematic reviews, conference proceedings and Masters or PhD theses will be excluded.

Information sources and search strategy
Searches will be conducted in six electronic databases: PubMed, CINAHL (via EBSCO), EMBASE (via Elsevier), CENTRAL (Cochrane Central Register of Controlled Trials; via Wiley Online Library) and Web of Science (via Clarivate Analytics), and for the grey literature, OpenGrey and Grey Literature Network Service. Additionally, hand searches of the reference lists of all included studies and previously published systematic reviews of MRI staging of PCa will be performed. The search strategy will be developed in consultation with a medical librarian with expertise in systematic review searching. The search terms will be adjusted to the specificities of the different databases. Keywords or database-specific subject headings (e.g., MeSH) and the Boolean operators ‘OR’ and ‘AND’ will be used to combine the search terms. The keywords included were “Prostate neoplasm”, “Machine learning”, “Artificial intelligence”, “Radiomics”, “Deep learning”, “Staging” and “Magnetic Resonance Imaging” (online supplemental file).

Study selection and data extraction
No area restriction will be applied and only studies published in the English language will be included. The search in each database will be performed from January 2007 to January 2021.

Two independent reviewers will conduct the selection process. All records identified in the search stage will be screened by title/abstract and studies clearly not matching the criteria will be discarded. The remaining studies will be full-text reviewed and included or discarded according to the inclusion/exclusion criteria. Any disagreement between the reviewers will be solved by consensus or by a third one if necessary. Reasons for the exclusion of full text records will be recorded. Details on the selection process of the studies will be documented into a flow chart following the PRISMA.

Assessment of risk of bias
Risk of bias of individual studies will be assessed independently by two reviewers. Since we will be including diverse types of studies, we will use different tools to assess the risk of bias depending of the characteristics of the studies. For all studies the RCT and non-randomised studies will be using the QUADAS-2 tool for assessing risk of bias. This tool covers seven sources of bias: (1) random sequence generation; (2) allocation concealment; (3) blinding of participants and personnel; (4) blinding of outcome assessment; (5) incomplete outcome data; (6) selective reporting; (7) other bias for RCT. For each one of the them the risk can be assessed as high risk, unclear risk or low risk depending of the information offered by the study. Non-randomised trials such as cohort studies or case control studies will be assessed based on three perspectives: (1) selection of study groups; (2) comparability of the groups; (3) ascertainment of exposure (in case–control studies) or outcome of interest (in cohort studies). Data from these studies will be extracted and tabulated, and then reviewed for risk of bias and applicability using the QUADAS-2 tool. All evaluations will be done in duplicate. Studies with a high risk of bias and low applicability will be excluded.

A narrative synthesis will be conducted, acknowledging the risk of bias and the strength and consistency of significant associations. We will extract and report all unadjusted and adjusted measures of association from included studies. Associations with outcome will be defined as a significant (p<0.05) univariable association, a significant (p<0.05) adjusted association (multivariable) or a significant (p<0.05) association in other predictive analysis (linear or multiple regression). Effect sizes will be represented as an OR or relative risk (RR) and considered as significant when the 95% CI do not include.

The results will be analysed using the levels of evidence proposed by Furlan et al.18: (A) strong evidence, defined as consistent (>75%) findings among multiple ≥2 high-quality studies; (B) moderate evidence: findings in one high-quality study and consistent (>75%) findings in ≥2 low-quality studies; (C) limited evidence: findings in one
high-quality study or consistent findings in 3 low-quality studies; (D) conflicting or inconclusive evidence: consistent findings in <75% of the studies, or results based on one single study.

The possibility of performing a formal meta-analysis will be evaluated, depending on the numbers, quality and outcome variables and effect measures.

**Patient and public involvement**

This is a protocol for a systematic review that will be based on previously published data, therefore no participant recruitment will take place. The involvement of participants on the recruitment and dissemination of results is not applicable.

**ETHICS AND DISSEMINATION**

This work will be based on data that is public and already published, therefore an ethical approval is not necessary. The result obtained from this work will be published in a peer-reviewed journal and disseminated in relevant conferences and thesis elaboration. If any amendments are needed due to deviations from this protocol in the execution of the study, they will be recorded and noted in the publication.

**Correction notice**

This article has been corrected since it was published Online First. One of the authors name has been updated.

**Contributors**

The manuscript protocol was drafted by AG and was revised by EN. NP contributed for the development of the selection criteria, the risk of bias assessment strategy and data extraction criteria. HD will develop the search strategy and manage the study records on Mendeley software. AG and EN will screen the studies obtained in the search for their eligibility criteria, extract data and assess risk of bias of included studies. NP will resolve any disagreements regarding eligibility for inclusion, data extraction and/or risk of bias. All authors have read, provided feedback and approved the final protocol. AG is the guarantor of this article.

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**Competing interests**

None declared.

**Patient and public involvement**

Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

**Patient consent for publication**

Not applicable.

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**Supplemental material**

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**REFERENCES**

Search strategy

The full search strategy for all databases is as follows:

- **Web of Science**: TOPIC (Machine learning OR Artificial intelligence OR Radiomics OR Deep learning) AND TOPIC: (prostate OR prostatic) AND TOPIC: (multiparametric magnetic resonance imaging OR mpMRI OR MRI OR magnetic resonance)
  
  Refined by: DOCUMENT TYPES: (ARTICLE OR DATA PAPER OR EARLY ACCESS OR CORRECTION OR RETRACTION)
  
  Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, ESCI, CCR-EXPANDED, IC
  
  Timespan=2000-2021

- **EMBASE**: The combination ('prostate cancer'/exp AND 'nuclear magnetic resonance imaging'/exp) AND ('artificial intelligence'/exp OR 'machine learning'/exp) OR ('machine learning':ab,ti OR 'artificial intelligence':ab,ti OR radiomics:ab,ti OR 'deep learning':ab,ti) AND (prostate:ab,ti OR mri:ab,ti OR 'multiparametric magnetic resonance imaging':ab,ti OR mpmri:ab,ti OR 'magnetic resonance':ab,ti)

  
  Filters applied: English, from 2000-3000/12/12.

- **CENTRAL**: "multiparametric magnetic resonance imaging OR mpMRI OR MRI OR magnetic resonance in Title Abstract Keyword AND Machine learning OR Artificial intelligence OR Radiomics OR Deep learning in Title Abstract Keyword AND prostatic neoplasms OR prostate OR prostatic in Title Abstract Keyword - with Publication Year from 2000 to 2021, with Cochrane Library publication date Between Jan 2000 and Feb 2021, in Trials (Word variations have been searched).
Search strategy

The full search strategy for all databases is as follows:

- **Web of Science**: TOPIC (Machine learning OR Artificial intelligence OR Radiomics OR Deep learning) AND TOPIC: (prostate OR prostatic) AND TOPIC: (multiparametric magnetic resonance imaging OR mpMRI OR MRI OR magnetic resonance)
  Refined by: DOCUMENT TYPES: (ARTICLE OR DATA PAPER OR EARLY ACCESS OR CORRECTION OR RETRACTION)
  Indexes=SCI-EXPANDED, SSCI, A&HCI, CPCI-S, CPCI-SSH, ESCI, CCR-EXPANDED, IC
  Timespan=2000-2021

- **EMBASE**: The combination ('prostate cancer'/exp AND 'nuclear magnetic resonance imaging'/exp) AND ('artificial intelligence'/exp OR 'machine learning'/exp) OR ('machine learning':ab,ti OR 'artificial intelligence':ab,ti OR radiomics:ab,ti OR 'deep learning':ab,ti) AND (prostate:ab,ti OR mri:ab,ti OR 'multiparametric magnetic resonance imaging':ab,ti OR mpmri:ab,ti OR 'magnetic resonance':ab,ti)


- **CENTRAL**: "multiparametric magnetic resonance imaging OR mpMRI OR MRI OR magnetic resonance in Title Abstract Keyword AND Machine learning OR Artificial intelligence OR Radiomics OR Deep learning in Title Abstract Keyword AND prostatic neoplasms OR prostate OR prostatic in Title Abstract Keyword - with Publication Year from 2000 to 2021, with Cochrane Library publication date Between Jan 2000 and Feb 2021, in Trials (Word variations have been searched).