

Multiparametric magnetic resonance imaging for pre-treatment local staging of prostate cancer: A Cancer Care Ontario clinical practice guideline

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Abstract

Introduction: The utility of T2-weighted magnetic resonance imaging (MRI) in the local staging of prostate cancer is controversial. Due to the success of multiparametric MRI in cancer localization, there is renewed interest in MRI (\pm functional sequences) for local staging. Guidance on pre-treatment local staging of prostate cancer by MRI was developed using systematic review methodology and expert consultation.

Methods: MEDLINE, EMBASE, Cochrane Database of Systematic Reviews, and other databases were searched to identify studies comparing: (1) MRI staging vs. radical prostatectomy staging on diagnostic accuracy outcomes; and (2) MRI staging vs. routine clinical staging on clinical and patient outcomes. Studies meeting inclusion criteria were synthesized by outcome and sensitivity/specificity analysis by tumour location was performed. Evidence quality of included studies was assessed and considered in recommendation formulation.

Results: The literature search identified 2510 citations; 62 studies were included. Analysis of MRI \geq 1.5 T plus endorectal coil (ER) (\pm functional sequences) in the detection of extraprostatic extension or seminal vesicle invasion showed modest sensitivities (\geq 50%) and excellent specificities ($>$ 85%) among patients scheduled for radical prostatectomy. MRI upstaging was shown in 20/21 studies, with large variation in correctness (11–85%). Scarcity of clinical and patient outcomes among studies limited synthesis and evaluation. Quality assessment found non-trivial biases.

Conclusions: Modest imaging performance was shown for MRI (1.5 T + ER and 3 T \pm ER) \pm functional sequences in regards to sensitivity. Limitations in study design, reporting of clinical and patient outcomes, and the heterogeneous use of MRI tempered the strength of the recommendations.

Introduction

Prostate cancer among men in Canada ranks first in terms of the number of new cases (21.6%) and third in terms of mortality (9.9%), after colorectal cancer (11.8%) and lung cancer (25.3%).¹

Prostate cancer is typically diagnosed through a combination of prostate-specific antigen (PSA) screening, digital rectal examination (DRE), and transrectal ultrasonography (TRUS)-guided biopsy. Limitations when locally staging prostate cancer for treatment planning purposes include: low specificity, missed regions of the prostate during DRE or biopsy (e.g., anterior tumours), and potentially inaccurate information regarding tumour volume, extent and aggressiveness of disease with DRE and non-targeted TRUS-guided biopsy.²

There is renewed interest in the benefits of magnetic resonance imaging (MRI) (\pm functional sequences) in the local staging of prostate cancer. T2-weighted MRI for local staging of prostate cancer traditionally involves high-resolution imaging, which delineates the prostatic anatomy in detail. MRI is non-invasive, can visualize the boundaries of the prostate, and with multiparametric techniques can also determine the location of clinically significant cancers.³ Multiparametric MRI (mpMRI) is the addition of two or more functional sequences to MRI, such as diffusion-weighted imaging (DWI), dynamic contrast-enhanced imaging (DCE), and proton magnetic resonance spectroscopic imaging (MRS). The current recommended combination of functional sequences to improve detection and localization of prostate cancer is MRI plus DWI and DCE. This form of mpMRI is gaining wide acceptance and is being adopted for image-guided biopsy, biopsy avoidance, and consideration of image-guided therapy of index tumours;⁴ however, MRI has not gained wide adoption for local staging of prostate cancer, as controversy still exists regarding its value in care and management.

Recognizing the potential significance of improved cancer localization by MRI and the need to more accurately stage patients compared to conventional methods to aid in radical treatment planning, the Cancer Imaging Program of Cancer Care Ontario (Toronto, Canada) in collaboration with the Prostate Cancer Disease Pathway Management Secretariat developed this clinical practice guideline. An expert group was assembled, including radiologists, radiation oncologists, urologists, and health research methodology as part of the Program in Evidence-Based Care (McMaster University, Canada) to answer the following research questions: what is the performance and diagnostic accuracy of MRI (\pm functional sequences including DWI, DCE, or MRS) in the pre-treatment local staging of prostate cancer and impact on clinical and patient outcomes?

Methods

A two-staged approach was used: review of existing relevant guidelines and a systematic review. Prior to finalization, the guideline underwent peer-review and stakeholder engagement.

Search strategy

Potentially relevant guidelines were identified and reviewed by searching online databases and guideline developer websites, and by performing a systematic literature search in standard databases. Search criteria included relevant publications in the last 10 years with clearly described methods and recommendations. MEDLINE, EMBASE, Cochrane Database of Systematic Reviews, and other databases were used to identify primary studies and systematic reviews (Appendix 1).

Study assessment

Potentially relevant studies were identified and reviewed on the basis of title and abstract by one reviewer (JS). Reference lists were also examined for additional relevant studies. Inclusion criteria were: (1) studies published between January 1, 2008 and February 17, 2016; (2) adults ≥ 18 years; (3) studies on the pre-treatment local staging of prostate cancer in men with newly diagnosed biopsy-confirmed prostate cancer and candidates for radical treatment; (4) studies of MRI ≥ 1.5 T \pm endorectal coil (ER) \pm DWI, DCE, MRS; (5) studies with at least one outcome of interest; and (6) minimum size of 30 patients. Studies were excluded if they investigated: (1) technical imaging aspects; (2) post-treatment or pre-diagnosis; (3) other combined technologies; and (4) active surveillance. Studies not in English, case reports, conference abstracts, and in vitro or animal studies were also excluded. There were no exclusions based on study

design. Outcomes of interest included diagnostic accuracy (e.g., sensitivity, specificity); clinical outcomes, including stage classification, risk stratification category, treatment plan, surgical margins, and biochemical recurrence; and patient outcomes (e.g., survival). Full-text articles of studies meeting inclusion criteria were retrieved. Potentially eligible systematic reviews were assessed for methodological quality,⁵ with a score ≥ 7 considered satisfactory. The quality of the evidence was appraised using standardized tools.⁶⁻⁸

Analysis

Data abstraction was performed by one abstractor (JS). Abstracted data included study variables, MRI variables, and outcomes. Scatterplots of sensitivity vs. 1-specificity for studies of MRI ≥ 1.5 T + ER \pm DWI, DCE, MRS were examined based on published estimates and stratified by tumour location (i.e., extraprostatic extension [EPE] vs. seminal vesicle invasion [SVI]). MRI studies of 1.5 T and 3 T without ER \pm DWI, DCE, MRS were also examined. Median sensitivities and specificities were calculated and interpreted.⁹ Change in stage and change in treatment were calculated from before and after MRI. The percentage of positive surgical margins was stratified by whether or not MRI results were used to inform the nature of radical prostatectomy surgery. Heterogeneity variables included field strength, ER, functional sequences, and sample size. All data were audited by an independent auditor. All analysis included studies of MRI \pm DWI, DCE, MRS.

Results

The literature search identified 2510 citations; 201 papers underwent full-text review. In total, there were 62 included studies (Fig. 1). One systematic review published as a component of a clinical practice guideline¹⁰ relevant to the objectives, questions, and outcomes of this guideline and with a satisfactory quality score helped to inform the literature search strategy and evidentiary base. Otherwise, a search for pre-existing guidelines did not yield an endorsable document. A summary of included primary studies is shown in Table 1.

By outcome, there were 48 primary studies and one systematic review/meta-analysis on diagnostic accuracy; 21 studies on stage classification; two studies on risk stratification; six studies on treatment plan; 19 observational studies and one randomized controlled trial on surgical margins; five studies on biochemical recurrence; and no studies with patient outcomes.

Recommendation 1

mpMRI use for pre-treatment local staging of prostate cancer is a reasonable option for assessment of EPE in intermediate-

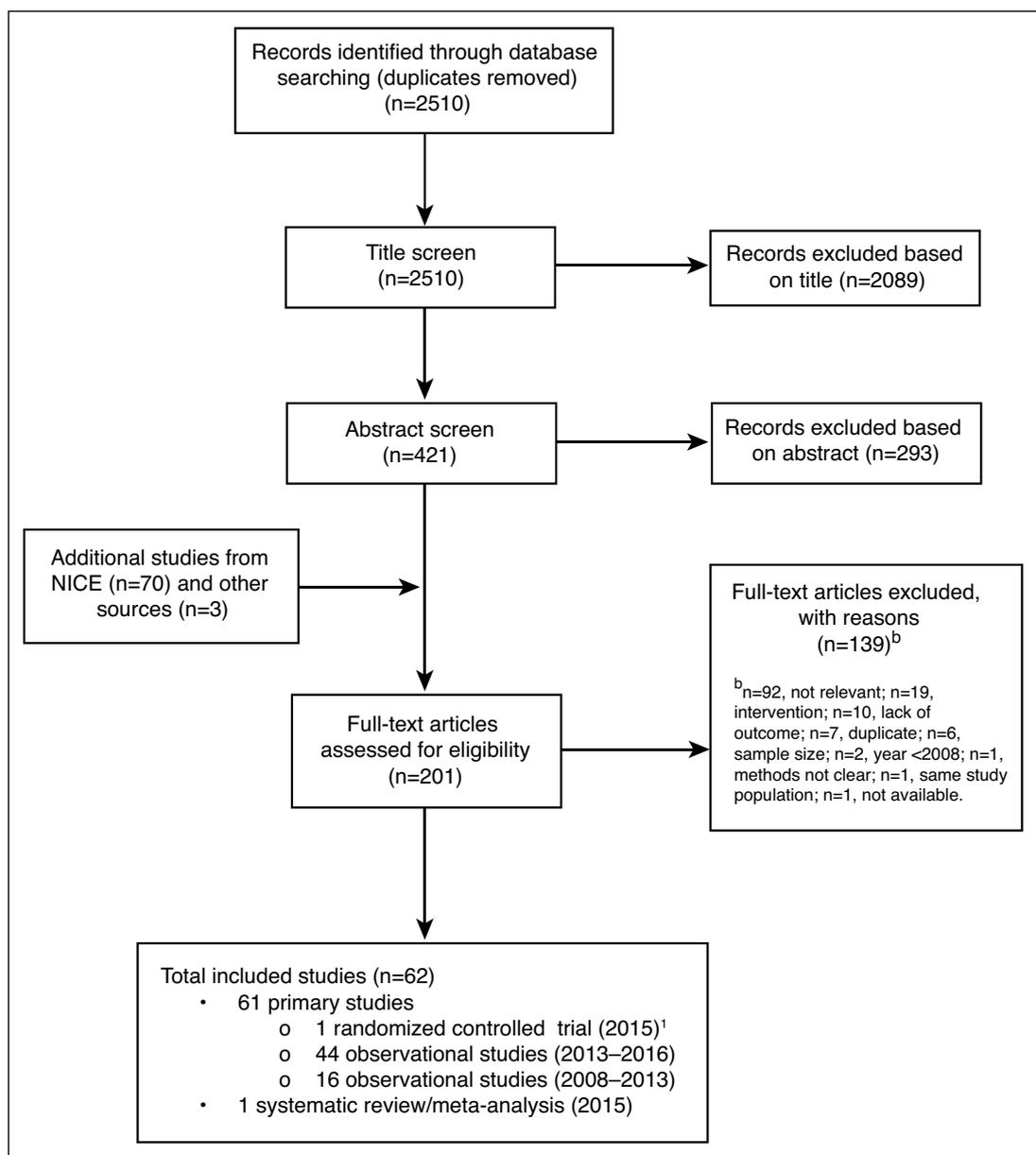


Fig. 1. Citation flow chart.¹ For stage classification, the prior study³⁶ upon which the more recent trial³³ was based was also used. NICE: National Institute of Clinical and Health Excellence.

and high-risk patients being considered for radical therapy if knowledge of EPE will alter management.

Key evidence

There were 19/49 (38.8%) primary studies of MRI ≥ 1.5 T + ER \pm DWI, DCE, MRS that considered the detection of EPE and/or SVI^{11–28} (Figs. 2, 3). Median sensitivities were modest and specificities were excellent, and similar to a recently published systematic review/meta-analysis²⁹ (Table 2).

There were 6/61 (9.8%) studies that reported on the outcome of change in treatment plan.^{30–35} All six studies were

consistent in showing increased intensity of therapy as a result of MRI staging, with <1–43% of patients experiencing increased therapy due to MRI staging.^{30–35} Three studies reported that MRI-informed treatment plans were correct, as shown in 63–97% of patients.^{31,32,35}

There were 21/61 (34.4%) studies that reported on the outcome of change in stage classification. There were 20/21 (95.2%) studies that consistently demonstrated upstaging by MRI,^{33,36} and upstaging by MRI compared to routine clinical staging.^{12,13,16,17,20,21,23,30,32,34,37–45} Upstaging by MRI was correct by pathology in seven studies, with a range from 11–85%.^{16,17,32,34,38,41,44}

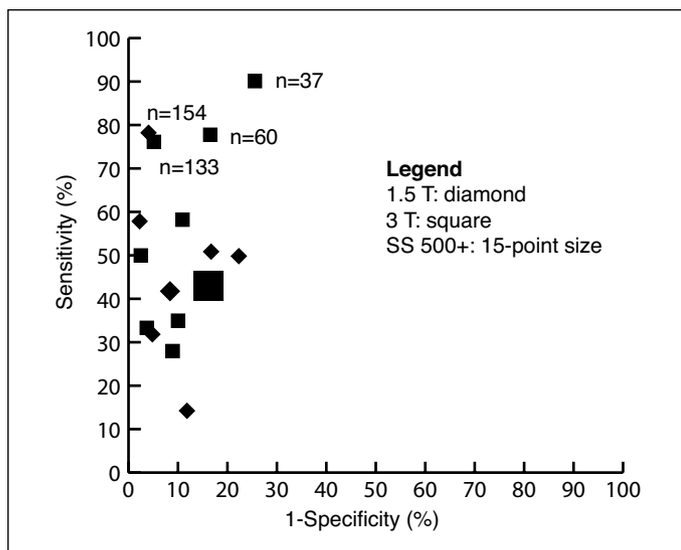


Fig. 2. Magnetic resonance imaging studies of ≥ 1.5 T + ER \pm DWI, DCE, MRS in EPE (n=17). DCE: dynamic contrast-enhanced imaging; DWI: diffusion-weighted imaging; EPE: extraprostatic extension; ER: endorectal coil; MRS: magnetic resonance spectroscopic imaging; SS: sample size; T: Tesla.

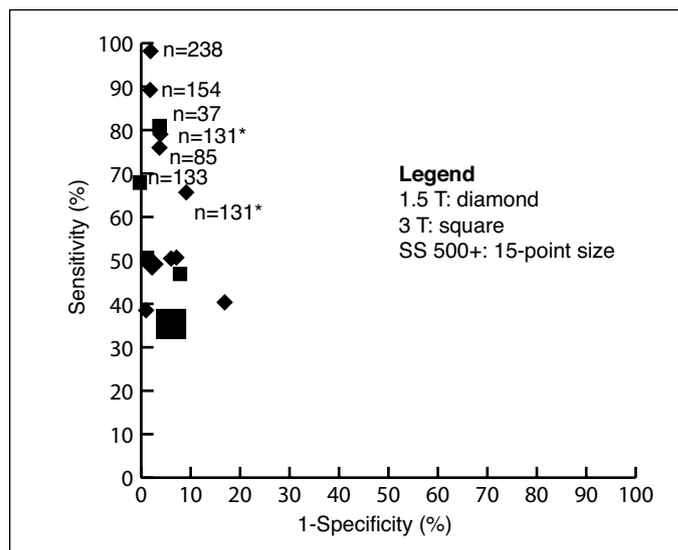


Fig. 3. Magnetic resonance imaging studies of ≥ 1.5 T + ER \pm DWI, DCE, MRS in SVI (n=15). *Denotes data point from same study population.²⁴ DCE: dynamic contrast-enhanced imaging; DWI: diffusion-weighted imaging; ER: endorectal coil; MRS: magnetic resonance spectroscopic imaging; SS: sample size; SVI: seminal vesicle invasion; T: Tesla.

Qualifying statements

The quality of evidence was judged to be of high risk. As a result, the above recommendation reflects a blended synthesis of evidence and expert opinion. mpMRI is best defined according to current standards.⁴ Furthermore, caution should be exercised when considering nerve-sparing surgery on the basis of mpMRI evaluation indicating no EPE on the side of prostate cancer.

Recommendation 2

Centres using mpMRI for local prostate cancer staging must have a quality assurance program in place to measure diagnostic performance.

Key evidence and qualifying statements

This recommendation is based on expert opinion with mpMRI best defined according to current standards.⁴ Quality assurance programs were not reviewed; however, use of standardized reporting has shown statistically significant improvements in sensitivity vs. non-standardized reporting.⁴⁶

Discussion

Systematic review methodology combined with expert consultation informed this clinical practice guideline. Based on observational evidence, modest imaging performance was shown for MRI (1.5 T + ER and 3 T \pm ER) \pm DWI, DCE, MRS in regards to sensitivity. Future high-quality diagnostic

accuracy studies and health outcomes studies are needed to inform evidence-based recommendations.

Our systematic review of the literature is consistent with a well-conducted, recently published systematic review/meta-analysis that reported overall sensitivity of approximately 50–60% and specificity of >85%, despite differences in methodology (e.g., years searched) and analyzed studies (e.g., ER, field strength, mpMRI).²⁹ In our work, above average sensitivities were achieved in some studies; however, whether this was due to the addition of functional sequences or field strength was not clear owing to too few studies (Table 3). Other reports of sensitivity and specificity of MRI for local staging of prostate cancer range from 15–100% and 67–100%, respectively.⁴⁷ Our synthesis of the published data showed a marked shift towards a higher minimum specificity value in the detection of EPE (74–98% vs. 49–99%, difference: 25%) and a shift towards a higher minimum sensitivity value in the detection of SVI (35–97% vs. 23–80%, difference: 12%).⁴⁷ In our work, higher performance was likely due to consideration of ER status. Among non-ER studies, 1.5 T had a lower sensitivity than 3 T (Table 2), which was also consistent with a previous detailed analysis.²⁹

The quality assessment of ER studies on imaging performance was judged to be of high risk. Explanations surround the lack of detail reported for radiology and pathology evaluation and interpretation, and the difficulty in distinguishing between microscopic to gross EPE.⁴⁸ Future studies should consider improvements in patient sampling and study design, standardized use of functional sequences, explicit pathology criteria, and blinding of both radiologists and pathologists.

Table 1. Summary of included primary studies (n=61)

Characteristic	Range or no. studies (%)
Age (median or mean)	58–70 years
Minimum time to imaging post-biopsy	2 weeks
Radiologist blinding	34 (55.7)
Field strength	
1.5 T	30 (49.1)
3 T	30 (49.1)
Both	1 (1.6)
Study type	
RCT	1 (1.6)
Retrospective	35 (57.3)
Prospective	19 (31.1)
Mixed	6 (9.8)
PSA level (median or mean)	
<10 ng/ml	41 (67.2)
10–20 ng/ml	19 (31.1)
>20 ng/ml	0 (0)
Not known	1 (1.6)
No ER use	
Plus body or pelvic coil	22 (36.1)
Minus body or pelvic coil	5 (8.2)
ER use	
Plus body or pelvic coil	27 (44.3)
Minus body or pelvic coil	7 (11.5)
Imaging	
MRI	17 (27.9)
MRI + DWI	10 (16.4)
MRI + (DCE and/or MRS)	6 (9.8)
MRI + DWI + (DCE and/or MRS)	28 (45.9)

DCE: dynamic contrast-enhanced imaging; DWI: diffusion-weighted imaging; ER: endorectal coil; MRI: magnetic resonance imaging; MRS: magnetic resonance spectroscopic imaging; PSA: prostate-specific antigen; RCT: randomized controlled trial.

The impact of pre-treatment local staging by MRI ± DWI, DCE, MRS on clinical and patient outcomes is still not clear. Although studies consistently demonstrated upstaging by imaging compared to clinical staging, this was only correct by pathology in a few studies and the magnitude was highly variable (11% to 85%),^{16,17,32,34,38,41,44} thus making any con-

clusions to be drawn imprecise. There was limited information on the outcome of risk stratification category,^{30,34} however, the results were consistent with the trend of upstaging. Changes to treatment plan with imaging were mixed, as they included both an increase³⁰⁻³⁵ and decrease^{31,35} to therapy intensity, with a tendency towards more intense therapy across included studies.³⁰⁻³⁵ The correctness of treatment changes was under-studied.^{31,32,35} The relation between tumour or EPE detected on imaging to biochemical recurrence was mixed and difficult to decipher due to unclear reporting and statistical methods.^{18,37, 42,43,49} The outcome of positive surgical margin status is a complex one, with its clinical impact increasingly scrutinized.⁵⁰ A recent randomized controlled trial did not show a beneficial effect of MRI+DWI³³ on surgical margins. There were a number of potential limitations to the trial that diluted the ability to detect a difference in surgical margin status between groups, including: limited power; specified criteria for deciding how to modify the surgical plan based on imaging (including a wider excision at sites of tumour) was not part of the study design; limitations of the surgical technique associated with robotic surgery, including the lack of a specified surgical protocol for various types of imaging findings; and the protocol detailing communication between the radiologist and the urologist could be improved.

The quality assessment of studies informing these outcomes was judged to involve non-trivial serious risk of biases. The limitations included small study sizes, paucity of data, the lack of consistently reported outcomes across studies, and the lack of comparable analysis methods across studies. Moreover, clinical and pathology differences between stratified groups being compared need statistical consideration.

Conclusion

In summary, modest imaging performance using MRI (1.5 T + ER and 3 T ± ER) ± DWI, DCE, MRS in the detection of EPE and SVI for patients scheduled to undergo radical prostatectomy was shown in regards to sensitivity. Our rec-

Table 2. Summary of diagnostic accuracy

Current guideline	No. studies	Sensitivity			No. studies	Specificity		
		Median	Min	Max		Median	Min	Max
ER use								
≥1.5 T for EPE	16	50.0	14.0	90.0	16	91.0	74.0	98.0
≥1.5 T for SVI	13	50.0	34.9	97.0	15	96.0	83.1	100.0
No ER use								
1.5 T (tumour, EPE, SVI)	9	36.2	0	81.3	9	90.3	65.0	97.7
3 T (tumour, EPE, SVI)	13	58.3	22.0	92.0	13	86.6	55.2	99.0
De Rooij et al (2015)	No. studies	Estimate	95% CI		No. studies	Estimate	95% CI	
EPE	45	0.57	0.49-0.64		45	0.91	0.88-0.93	
SVI	34	0.58	0.47-0.68		34	0.96	0.95-0.97	
Stage T3	38	0.61	0.54-0.67		38	0.88	0.85-0.91	

CI: confidence interval; EPE: extraprostatic extension; ER: endorectal coil; SVI: seminal vesicle invasion; T: Testa.

Table 3. Characteristics of diagnostic accuracy studies (+ER)

Characteristic	No. data or studies (%)
EPE	
Field strength	
1.5 T	9 (52.9)
3 T	8 (47.1)
Functional techniques	
MRI	8 (47.1)
MRI + DWI	1 (5.9)
MRI + (DCE and/or MRS)	0 (0)
MRI + DWI + (DCE and/or MRS)	8 (47.1)
Studies of >60% sensitivity	
1.5 T	1 (25.0)
3 T	3 (75.0)
MRI	1 (25.0)
MRI + DWI	0 (0)
MRI + (DCE and/or MRS)	0 (0)
MRI + DWI + (DCE and/or MRS)	3 (75.0)
SVI	
Field strength	
1.5 T	10 (66.7)
3 T	5 (33.3)
Functional techniques	
MRI	9 (60.0)
MRI + DWI	1 (6.7)
MRI + (DCE and/or MRS)	0 (0)
MRI + DWI + (DCE and/or MRS)	5 (33.3)
Studies of >60% sensitivity	
1.5 T	5 (71.4)
3 T	2 (28.6)
MRI	3 (42.9)
MRI + DWI	0 (0)
MRI + (DCE and/or MRS)	0 (0)
MRI + DWI + (DCE and/or MRS)	4 (57.1)

DCE: dynamic contrast-enhanced imaging; DWI: diffusion-weighted imaging; EPE: extraprostatic extension; ER: endorectal coil; MRI: magnetic resonance imaging; MRS: magnetic resonance spectroscopic imaging; SVI: seminal vesicle invasion; T: Tesla.

ommendations were tempered due to limited evidence. The recommendations are best used in the context of the current Cancer Care Ontario Prostate Cancer Diagnosis Pathway⁵¹ and mpMRI standards.⁴

Competing interests: Dr. Morash has been an Advisor for Abbvie, Astellas, AstraZeneca, Ferring, Janssen, and Sanofi. Dr. Morgan has been an Advisor for Abbvie, Accuray, Amgen, Astellas, Bayer, Janssen, and Sanofi; and has participated in clinical trials for Bayer and Janssen. Dr. Haider has been an Advisor for Bayer and has presented at numerous scientific meetings about MRI and advocates for its use in planning and staging in prostate cancer. All remaining authors report no competing personal or financial interests.

Members of the Guideline Development Group also disclosed potential conflicts of interest. One group member receives funding and is a trial co-investigator for MRI and prostate cancer and has published on MRI and radiotherapy planning. Another member receives support from an imaging company and has grant application activity in prostate imaging.

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Appendix 1. Supplemental methods: Literature search strategy

Using the following databases: Cochrane Central Register of Controlled Trials, Cochrane Database of Systematic Reviews, Database of Abstracts of Reviews of Effects, Health Technology Assessment, Ovid MEDLINE In-Process & Other Non-Indexed Citations, Ovid MEDLINE, EMBASE

- 1 exp Prostatic Neoplasms/
- 2 (prostat\$ adj3 (cancer\$ or carcinoma\$ or adeno\$ or malignan\$ or tumo?r\$ or neoplas\$)).mp.
- 3 1 or 2
- 4 exp Neoplasm Staging/
- 5 (staging or stage\$1 or classif\$ or evaluat\$ or tnm).mp.
- 6 4 or 5
- 7 exp Magnetic Resonance Imaging/
- 8 Magnet\$ resonance.mp.
- 9 (MRI or MR\$2 or NMR\$1).mp.
- 10 (MR adj (imag\$ or scan\$)).mp.
- 11 (magnet\$ adj (imag\$ or scan\$)).mp.
- 12 ((magnet\$ or MR) adj spectroscop\$).mp.
- 13 or/7-12
- 14 (magnetic resonance imag\$ or magnetic resonance spectroscop\$).mp.
- 15 (dynamic adj4 (MRI or magnet\$)).mp.
- 16 (diffusion weight\$ adj3 (MRI or magnet\$)).mp.
- 17 ((T1-weighted or T2-weighted or T3-weighted) adj3 imag\$).mp.
- 18 (magnet\$ adj (imag\$ or spectroscop\$ or scan\$ or resonance)).mp.
- 19 (MPMRI or MP-MRI or MR\$2 or DWI\$ or DW-MRI or DCE\$ or fmri).mp.
- 20 or/14-19
- 21 13 or 20
- 22 (comment or letter or editorial or note or erratum or short survey or news or newspaper article or patient education handout or case report or historical article).pt.
- 23 exp animal/ not humans/
- 24 22 or 23
- 25 (3 and 6 and 21) not 24
- 26 limit 25 to (english language and yr="2013-Current") [Limit not valid in CDSR,DARE; records were retained]
- 27 remove duplicates from 26