Synthetic magnetic resonance imaging for primary prostate cancer evaluation: Diagnostic potential of a non-contrast-enhanced bi-parametric approach enhanced with relaxometry measurements

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HIGHLIGHTS
• Diagnostic performances of synthetic bpMRI and conventional bpMRI are comparable for primary PCa
• Diagnostic performance of synthetic MRI variables are similar to that of DCE-MRI for csPCa in PZ
• Synthetic bpMRI shows potential as a contrast agent-free method for primary PCa

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ABSTRACT

Purpose: Bi-parametric magnetic resonance imaging (bpMRI) with diffusion-weighted images has wide utility in diagnosing clinically significant prostate cancer (csPCa). However, bpMRI yields more false-negatives for PI-RADS category 3 lesions than multiparametric (mp)MRI with dynamic-contrast-enhanced (DCE)-MRI. We investigated the utility of synthetic MRI with relaxometry maps for bpMRI-based diagnosis of csPCa.

Methods: One hundred and five treatment-naïve patients who underwent mpMRI and synthetic MRI before prostate biopsy for suspected PCa between August 2019 and December 2020 were prospectively included. Three experts and three basic prostate radiologists evaluated the diagnostic performance of conventional bpMRI and synthetic bpMRI for csPCa. PI-RADS version 2.1 category 3 lesions were identified by consensus, and relaxometry measurements (T1-value, T2-value, and proton density [PD]) were performed. The diagnostic performance of relaxometry measurements for PI-RADS category 3 lesions in peripheral zone was compared with that of DCE-MRI. Histopathological evaluation results were used as the reference standard. Statistical analysis was performed using the areas under the receiver operating characteristic curve (AUC) and McNemar test.

Results: In 102 patients without significant MRI artefacts, the diagnostic performance of conventional bpMRI was not significantly different from that of synthetic bpMRI for all readers (p = 0.11–0.79). The AUCs of the combination of T1-value, T2-value, and PD (T1 + T2 + PD) for csPCa in peripheral zone for PI-RADS category 3 lesions were compared with that of DCE-MRI. Histopathological evaluation results were used as the reference standard. Statistical analysis was performed using the areas under the receiver operating characteristic curve (AUC) and McNemar test.

Abbreviations: ADC, apparent diffusion coefficient; BpMRI, Bi-parametric magnetic resonance imaging; csPCa, clinically significant prostate cancer; DCE-MRI, dynamic contrast-enhanced MRI; DWI, diffusion-weighted imaging; ISUP, International Society of Urological Pathology; MpMRI, Multiparametric MRI; MRI, magnetic resonance imaging; PZ, prostate cancer; PD, proton density; PI-RADS v2.1, Prostate Imaging Reporting and Data System version 2.1; PZ, peripheral zone; ROI, region of interest; TE, echo time; TR, repetition time; TZ, transition zone; T2WI, T2-weighted imaging.

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1. Introduction

Prostate cancer (PCa) is the most common cancer and the second leading cause of cancer-related deaths in men [1]. Although the life expectancy for localised PCa has markedly increased over the past few decades, distinguishing clinically significant PCa (csPCa) by imaging is challenging due to the multi-focal nature of the disease [2,3]. Magnetic resonance imaging (MRI) has been widely applied for PCa detection [4,5], and multiparametric MRI (mpMRI) (a combination of T2-weighted imaging [T2WI], diffusion-weighted imaging [DWI], and dynamic contrast-enhanced MRI [DCE-MRI]) has been established for csPCa detection and prediction of patient outcomes [6,7]. The important role of mpMRI has also been acknowledged in the Prostate Imaging and Reporting and Data System version 2.1 (PI-RADS v2.1) [8,9]. DWI has been used as an essential functional sequence for peripheral zone (PZ) PCa, while T2WI is considered the leading sequence for transition zone (TZ) PCa. For the PZ, if the DCE-MRI findings meet the criteria, lesions in category 3 can be reclassified to category 4 [8,9].

Recently, bi-parametric (bp) MRI (a combination of T2WI and DWI) has been proposed as an alternative to mpMRI with gadolinium-enhanced sequences, reducing the costs and potential side effects of gadolinium-based contrast agents [9–11]. According to a meta-analysis based on the PI-RADS v2 criteria, both methods achieve a comparable diagnostic performance for csPCa [12]. However, recent studies have indicated that the diagnostic performance of bpMRI is inferior to that of mpMRI when interpreted by non-expert radiologists [13,14]; moreover, the sensitivity of bpMRI for csPCa is lower than that of mpMRI when the PI-RADS v2.1 system is used, particularly for category 3 lesions [9,15]. Therefore, when evaluating csPCa without contrast media in bpMRI, the arising challenge is to improve the diagnostic accuracy of lesions that are determined to be category 3 in bpMRI but would be reclassified to category 4 when relying on DCE-MRI findings in an mpMRI protocol.

Synthetic MRI is an emerging technique that synthesises MR images at arbitrary contrast after the actual MR acquisition, which may allow for the quantitative assessment of lesions without contrast media [16–20]. In a recent study, Cui, et al. revealed that the diagnostic performance of relaxometry maps (T1-value, T2-value, and proton density [PD]) was inferior to that of the apparent diffusion coefficient (ADC) values for discriminating between PCa and other benign entities [21]. However, comparison of the diagnostic performances between relaxometry maps and DCE-MRI-based PI-RADS scores remains unknown.

The purpose of this study was to investigate the utility of synthetic MRI for bpMRI evaluation of primary PCa by assessing the impact of the reader’s experience (expert or basic prostate radiologists) on the bpMRI-based diagnostic performance and comparing the diagnostic performance of relaxometry maps derived from synthetic MRI with that of DCE-MRI for category 3 lesions in PZ.

2. Materials and methods

This prospective study was approved by the medical ethics committee of our institution (approval number: 20190148), and written informed consent was obtained from all patients included.

2.1. Patients

One hundred and five treatment-naïve patients suspected to have PCa based on an elevated serum prostate-specific antigen level or abnormal digital rectal examination, who underwent conventional prostate mpMRI, including T2WI, DWI, and DCE-MRI, prior to MRI-ultrasonography fusion-guided prostate-targeted biopsy (MRGB) between August 2019 and December 2020, were prospectively included in this study. All patients also underwent synthetic MRI at the time of conventional prostate mpMRI. Patients whose MRI scans were degraded due to prominent susceptibility artefacts from previous hip replacement (n = 3) were excluded. The remaining 102 patients were eligible for analysis (Fig. 1).

2.2. Image acquisition

Conventional prostate mpMRI and synthetic MRI were acquired using a 3.0-T system (DISCOVERY MR750 • SIGNA Pioneer; GE Healthcare, WI, USA) (Fig. 2). The parameters for DWI comprised: repetition time (TR)/echo time (TE), 4500/55.3 ms; and two different b-values (50 and 800 s/mm²). Computed intermediate and high b-value DWI (1000, 1500, and 2000 s/mm²) and ADC maps were generated from the native DWI dataset (i.e., 50 and 800 s/mm²) on the MR console. The parameters used for T2WI comprised: TR/TE, 6000/120 ms; DCE-MRI: TR/TE, 4/1.1, 2.2 ms. Data acquisition for DCE-MRI began simultaneously with the start of intravenous injection of a gadolinium-based contrast medium at 0.1 mmol/kg of gadobutrol (Gadovist, Bayer Schering Pharma) at a rate of 1.5 mL/s via power injector, followed by a 30-mL saline flush at the same rate as contrast medium injection. Multiphase DCE-MR images were obtained every 7 s for 210 s (30 phases). The parameters for synthetic MRI comprised: TR/TE, 4500/13.6, 88.5 ms [16]. Synthetic T2WI was generated for TR/TE, 6000/100 ms on the MR console prior to the reading evaluation. The sequence parameters used for conventional mpMRI and synthetic MRI are shown in Supplementary Table 1.

2.3. Image analysis

2.3.1. Subjective assessment of image quality

Two bpMRI datasets were sent to our institutional imaging server system for review.

![Patient selection flowchart](image-url)

**Patient selection**

**Comparison of synthetic bpMRI and conventional bpMRI**

**Comparison of relaxometry maps and DCE-MRI PI-RADS score in PZ**

**Conclusion:** Synthetic MRI with relaxometry maps shows promise for contrast media-free evaluation of csPCa.
Both bpMRI datasets were reviewed along with the ADC maps for reference, using a multimodality workstation (Centricity™ Universal Viewer, PACS, GE Healthcare, WI, USA). The reviewers were blinded to the type of sequence and parameters under evaluation. The two bpMRI datasets were independently evaluated by six radiologists (three experts who had performed prostate MR evaluations in >1000 cases and three basic prostate radiologists who had performed prostate MR evaluations in >400 cases based on the criteria of the ESUR/ESUI consensus statements) [13], who were blinded to all clinical and pathological data, including any previous imaging data. The two bpMRI datasets of each patient were assigned anonymous identifiers and reviewed in a random order [22,23]. For each reader, there was at least a 1-month interval between reading pairs of images (i.e., Conventional bpMRI and Synthetic bpMRI) to avoid recall bias.

While reviewing each T2WI scan (synthetic T2WI and conventional T2WI), all six readers subjectively evaluated each image on a scale of 1–5 (with a score of 1 corresponding to the worst quality and a score of 5 corresponding to the highest quality) for the following items: contraction artefacts (i.e., artefacts resulting from bowel peristalsis and contraction of the urinary bladder), motion artefacts (i.e., breathing and patient movement), and overall image quality (the image quality was evaluated per T2W-series) [24,25].

2.4. PI-RADS scoring evaluation

For each of the four prostate regions, i.e., right and left TZ, and right and left PZ, DWI, and T2WI scans of each bpMRI dataset were scored using a five-point scoring system, with DWI as the dominant sequence for the PZ and T2WI for the TZ, to obtain an overall score according to the PI-RADS v2.1 criterion [9]. A PI-RADS score between 1 (low likelihood) and 5 (high likelihood) was assigned to denote csPCa (International Society of Urological Pathology (ISUP) grade group ≥ 2 tumours) [9]. When multiple tumours were present in one prostate region (i.e., right and left TZ, and right and left PZ), the tumour with the highest score (index lesion) was used to calculate the score for that prostate region.

2.5. Comparison between relaxometry maps derived from synthetic MRI and DCE-MRI PI-RADS scores for category 3 lesions in PZ

Subsequently, three expert prostate radiologists and three basic prostate radiologists each identified all PI-RADS category 3 lesions on synthetic bpMRI in the PZ by consensus and placed a region of interest (ROI) on the lesion to measure the relaxometry values of the lesion. Dedicated analysis software for synthetic MRI automatically calculated the relaxometry values. The mean T1, T2, and PD values of the pixels within the ROI were used as relaxometry measurements for further analyses. For groupwise comparisons, the mean of the values measured by each of the three readers was used for statistical analysis [26]. The PD level of pure water was set at 100%.

At the time of relaxometry map analyses, the DCE-MRI PI-RADS score (positive/negative) was determined by both expert and basic radiologists in each case by consensus based on the PI-RADS v2.1 criteria [9]. Lesions featuring focal enhancements occurring earlier or concurrent to the enhancement of contiguous normal prostate tissue, which generally corresponds to a suspicious finding in T2WI and/or DWI, were assigned a positive DCE-MRI PI-RADS score [9].

2.6. Reference standard

The radical prostatectomy specimen was selected as the final reference.
proposed as an alternative to mpMRI with gadolinium-enhanced sequences. However, it has some disadvantages, including high costs and potential side effects from exposure to gadolinium. mpMRI is widely employed for PCa evaluation, particularly for the diagnosis of csPCa prior to prostate biopsy; however, it has some disadvantages, including high costs and potential side effects from exposure to gadolinium.

The clinicopathological data are shown in Table 1. The median time between MRI and subsequent prostate biopsies was 4 weeks (range, 1–6 weeks). All 102 included patients received MRGB. A total of 52 patients were diagnosed with PCa, and the number of regions diagnosed with PCa was 1/2/3/4 in 18/22/8/4 patients. Among the 52 PCa patients, 37 underwent radical prostatectomy.

Pca was present in 102 regions (59/43 in TZ/PZ) out of a total of 408 regions (204/204 in TZ/PZ); ISUP grade group 1 (n = 8/6), grade group 2 (n = 22/16), grade group 3 (n = 16/11), grade group 4 (n = 11/9), and grade group 5 (n = 2/1). The non-cancerous lesions (n = 145/161) consisted of benign prostate hyperplasia, prostatitis, and normal TZ or PZ.

Recently, bpMRI has been proposed as an alternative to mpMRI with gadolinium-enhanced sequences [12]. Similarly, the need for reducing false-negatives in PI-RADS category 3 lesions and improving the assessment of category 3 lesions without contrast administration remain important challenges [9, 13–15].

3.1. Theory

MpmMRI has been widely employed for PCa evaluation, particularly for the diagnosis of csPCa prior to prostate biopsy; however, it has some disadvantages, including high costs and potential side effects from gadolinium-based contrast agents [9–11]. Recently, bpMRI has been proposed as an alternative to mpMRI with gadolinium-enhanced sequences [12]. Similarly, the need for reducing false-negatives in PI-RADS category 3 lesions and improving the assessment of category 3 lesions without contrast administration remain important challenges [9, 13–15].

3.2. Calculation

Synthetic MRI would be beneficial as a non-contrast-enhanced protocol for the diagnosis of csPCa, if the diagnostic performance of synthetic bpMRI was comparable to that of conventional bpMRI and its relaxometry maps, which may represent the signature of csPCa [20,21], and provide a diagnostic performance similar to the DCE-MRI-based PI-RADS score for category 3 lesions in PZ.

4. Results

4.1. Patients’ and PCa lesions’ characteristics

The clinicopathological data are shown in Table 1. The median time between MRI and subsequent prostate biopsies was 4 weeks (range, 1–6 weeks). All 102 included patients received MRGB. A total of 52 patients were diagnosed with PCa, and the number of regions diagnosed with PCa was 1/2/3/4 in 18/22/8/4 patients. Among the 52 PCa patients, 37 underwent radical prostatectomy.

Pca was present in 102 regions (59/43 in TZ/PZ) out of a total of 408 regions (204/204 in TZ/PZ); ISUP grade group 1 (n = 8/6), grade group 2 (n = 22/16), grade group 3 (n = 16/11), grade group 4 (n = 11/9), and grade group 5 (n = 2/1). The non-cancerous lesions (n = 145/161) consisted of benign prostate hyperplasia, prostatitis, and normal TZ or PZ.

4.2. Comparison of the subjective image quality scores

The subjective image quality grading (average score) of each T2WI dataset (conventional vs. synthetic) of the 102 included patients is shown in Supplementary Table 2. No significant difference was observed between the two datasets regarding contraction artifacts, motion artifacts and overall image quality for both TZ and PZ, for all the six readers (p = 0.27–0.81). Representative cases of subjective scores of overall image quality are shown in Supplementary Fig. 1.

4.3. Comparison of the conventional and synthetic bpMRI datasets for primary PCa evaluation according to the PI-RADS criteria

The diagnostic performance of the two bpMRI datasets for csPCa is shown in Table 2. The Fleiss’ kappa values for interobserver agreement regarding PI-RADS v 2.1 overall score among all six radiologists were 0.76 and 0.79 for the TZ and PZ, respectively, in conventional bpMRI, and 0.75 and 0.78 for the TZ and PZ, respectively, in synthetic bpMRI. The Fleiss’ kappa values for interobserver agreements regarding PI-RADS v 2.1 overall score among expert/basic prostate radiologists were 0.82/0.78 and 0.80/0.78 for the TZ and PZ, respectively, in conventional bpMRI, and 0.79/0.78 and 0.83/0.82 for the TZ and PZ, respectively, in synthetic bpMRI.

For the TZ, the AUCs for the diagnosis of csPCa using conventional bpMRI/synthetic bpMRI evaluated by expert prostate radiologists 1, 2, 38 (64.4)/ 27 (62.8)a

Table 1

| Characteristics of the patients (n = 52) and their index PCa lesions (n = 102). |
|-----------------------------|--------------------|-----------------------------|-----------------------------|
| Patient characteristics   |                    |                             |                             |
| Age, years                  | 67.7 (44–85)a       | PSA level, ng/mL            | 7.82 (4.25–18.3)a           |
| Prostate cancer characteristics |                   |                               |                             |
| Clinical stage              | 2 (3.9)/29 (55.8)/5 (9.6)/10 (19.2)/5 (9.6)/1 (1.9)b | Site of the index lesion      | 29 (28.4)/30 (29.4)/22 (21.6)/21 (20.6)b |
| (T1c/T2a/T2b/T2c/T2a/T3b)   |                   | Maximum lesion length, mm    | 7.4 (0.25–7.1)/ 7.1 (0–22.9)b |
| Dominant lesion Gleason score (T2/PZ) |                 |                             |                             |
| 6                           | 8 (13.6)/ 6 (14.0)a | 7                           | 38 (64.4)/ 27 (62.8)a       |
| 8–10                        | 13 (22.0)/ 10 (23.2)b |                             |                             |
| PCa: prostate cancer; PSA: prostate-specific antigen; Rt: right; Lt: left; T2: transition zone; PZ: peripheral zone | |                             |                             |
| a Data presented in terms of the median (range) | | b Data presented in terms of the number (percentage) | |
T2-weighted imaging (T2WI) scans; Synthetic bpMRI: axial DWI scans with a b-value of 2000 s/mm²

...no significant difference in diagnostic performance between conventional bpMRI and synthetic bpMRI for all readers (p = 0.46, 0.21, and 0.49, for each reader). Furthermore, there was no significant difference in sensitivity, specificity, positive predictive value, and negative predictive value between conventional bpMRI and synthetic bpMRI for both TZ and PZ between the six readers (p = 0.11–0.78).

4.4. Synthetic MRI-based relaxometry measurements for category 3 lesions in PZ

Among the 41 lesions in PZ that were classified by the expert prostate radiologists as PI-RADS category 3, 12 (29.3%), 2 (4.9%), and 27 (65.8%) were pathologically determined to be csPCa, clinically insignificant PCa, and non-cancerous, respectively. Similarly, among the 41 lesions in PZ that were classified by the basic prostate radiologists as PI-RADS category 3, 12 (29.3%), 2 (4.9%), and 27 (65.8%) were pathologically determined as csPCa, clinically insignificant PCa, and non-cancerous, respectively. The
Fleiss’ kappa values for interobserver agreements regarding relaxometry measurements among expert and basic prostate radiologists were 0.89 and 0.87, respectively. The distribution of relaxometry measurements for csPca in PZ is shown in Fig. 3. For both expert and basic prostate radiologists, T1-value, T2-value, and PD of csPca were significantly lower than that of other entities in the PZ (p < 0.05). Overall, the combination of T1-value, T2-value, and PD (T1 + T2 + PD) had the highest AUC among all the combinations of relaxometry measurements for csPca for the PZ: 0.85 for expert prostate radiologists and 0.86 for basic prostate radiologists, respectively.

4.5. Comparison between relaxometry maps derived from synthetic MRI and DCE-MRI PI-RADS scores for category 3 lesions

The sensitivity and specificity of relaxometry measurements from synthetic MRI and DCE-MRI PI-RADS scores for the diagnosis of csPca in category 3 lesions in PZ are shown in Table 3. For the PZ, the AUCs of T1 + T2 + PD/DCE-MRI PI-RADS score for the diagnosis of csPca in category 3 lesions were 0.85/0.83 for expert radiologists and 0.86/0.83 for basic radiologists, with no significant difference between T1 + T2 + PD vs. DCE-MRI PI-RADS score for both groups of readers (p = 0.45, and 0.29, respectively). Furthermore, there was no significant difference in sensitivity and specificity between T1 + T2 + PD vs. DCE-MRI PI-RADS score for the PZ between the expert and basic groups (p = 0.35–1.00).

| Prostate Radiologist | Parameter | Relaxometry maps from synthetic MRI | DCE-MRI PI-RADS score | p-value |
|----------------------|-----------|-------------------------------------|------------------------|---------|
| Expert               | Sensitivity, % | 83.3 | 83.3 | 1.00 |
|                      | Specificity, % | 79.3 | 82.8 | 0.48 |
| Basic                | Sensitivity, % | 83.3 | 83.3 | 1.00 |
|                      | Specificity, % | 78.8 | 81.8 | 0.35 |

Table 3

Comparison of the diagnostic performance for csPca diagnosis between relaxometry measurements from synthetic MRI and the DCE-MRI PI-RADS score for category 3 lesions in PZ according to PI-RADS version 2.1.

The sensitivity and specificity of relaxometry maps from synthetic MRI and DCE-MRI PI-RADS scores for the diagnosis of csPca in category 3 lesions in PZ, for both expert and basic prostate radiologists. Therefore, the potential weaknesses of bpMRI evaluation, where the diagnostic performance is inferior to that of mpMRI for non-expert radiologists [13,14], and the sensitivity values of bpMRI are lower than those of mpMRI when using the PI-RADS v2.1 system for category 3 lesions [9,15], may be mitigated by combining (synthetic) bpMRI with relaxometry maps. As such, synthetic MRI has a potential as a non-contrast-enhanced method for primary PCa evaluation.

5. Discussion

We investigated the feasibility of bpMRI with relaxometry maps using synthetic MRI for primary PCa evaluation. We demonstrated that the diagnostic performance of synthetic bpMRI was comparable to that of conventional bpMRI for primary PCa evaluation in both TZ and PZ, for both expert and basic prostate radiologists. Additionally, the diagnostic performance of bpMRI observed in this study showed a comparable though slightly better diagnostic performance than that was reported in a recent meta-analysis by Cuocolo, et al. (AUC, 0.84) [30] and matches the results of a recent meta-analysis by Woo, et al. on the diagnostic accuracy of mpMRI for PCa (AUC, 0.90) [12]. Furthermore, the sensitivity and specificity of relaxometry maps from synthetic MRI were similar to those of the DCE-MRI PI-RADS scores for the diagnosis of csPca in PI-RADS category 3 lesions in the PZ, for both expert and basic prostate radiologists. Therefore, the potential weaknesses of bpMRI evaluation, where the diagnostic performance is inferior to that of mpMRI for non-expert radiologists [13,14], and the sensitivity values of bpMRI are lower than those of mpMRI when using the PI-RADS v2.1 system for category 3 lesions [9,15], may be mitigated by combining (synthetic) bpMRI with relaxometry maps. As such, synthetic MRI has a potential as a non-contrast-enhanced method for primary PCa evaluation.

Synthetic MRI, which is an emerging MRI acquisition technique, requires no additional image acquisition time for generating any arbitrary image contrasts after MRI acquisition, and does not require the use...
of any contrast agents [15–18]. According to a recent report by Hagiwara et al., double inversion recovery images without contrast agents derived from synthetic MRI enabled detection of more plaques in multiple sclerosis patients than conventional double inversion recovery images from dedicated brain MRI within a comparable acquisition time [31]. Moreover, according to a recent report by Yi et al., conventional knee MRI and synthetic MRI showed a substantial-to-almost perfect degree of agreement for the assessment of internal derangement of knee joints [32]. In the image quality analysis of our study, all subjective quality measures analysed (contraction artefacts, motion artefacts, and overall image quality) were comparable between synthetic bpMRI and conventional bpMRI. As indicated in previous studies, the diagnostic performance of bpMRI is inferior to that of mpMRI when interpreted by non-expert radiologists [13,14]. However, as shown in the results of this study, the diagnostic performance of synthetic bpMRI for basic prostate radiologists was similar to that for expert prostate radiologists for primary PCa diagnosis. Therefore, incorporating synthetic MRI into standard clinical PCa screening protocols may be useful, even for basic prostate radiologists.

Previously published literature about relaxometry MRI methods particularly focused on the T2-value for tissue characterisation and has shown that the T2-value of PCa, which has been recognised as a quantitative biomarker reflecting the mobile water content in different tissues, was significantly lower in csPCa than in non-cancerous lesions [33]. However, according to a recent report by Panda et al., the T1-value for category 3 csPCa is also lower than that for non-cancer in the T2 [26]. Our results confirm that both T1-value and T2-value, as quantitatively measured using synthetic MRI, are useful for the discrimination of csPCa from other (benign) entities in PZ. Moreover, as shown in the results of our study, T1 + T2 + PD yielded the highest AUCs. According to the report by Panda et al., the AUC of the combination of T1-value and ADC was 0.81 [26]. Our results (based on T1 + T2 + PD with AUCs of 0.85–0.91) were slightly better than the diagnostic performance metrics published previously.

From a clinical point of view, there are several potential advantages of using synthetic MRI for PCa diagnosis. First, synthetic MRI is a potentially time-saving method due to its relatively shorter examination time; however, if the scan fails, the entire scan has to be repeated, instead of only repeating a single sequence as in conventional MRI [10, 16–18]. However, the latter did not occur in any of the patients in our study. It also avoids exposure of patients to the potential risks of gadolinium-based contrast agents (i.e., nephrogenic systemic fibrosis and gadolinium deposition in the brain) [34–36]. Synthetic MRI also nullifies the risk of allergic reactions to contrast agents.

5.1. Limitations

Our study has several limitations. First, we divided the prostate into four regions, and only the index lesion with the highest pathological grade group or size was analysed in patients with multiple lesions, which might have resulted in selection bias. Nevertheless, the concept of the index lesion has been widely used in previous studies, including studies on PI-RADS [8,9,23,24]. Second, MRGB was used as the reference standard in several patients. Histological–radiological correlation is a great challenge in prostate MRI research. In this study, the radical prostatectomy result was used as the final pathological reference standard to minimise histological–radiological mismatch in patients who underwent both MRGB and subsequent radical prostatectomy. The concept of this type of reference standard (radical prostatectomy or MRGB) has been widely used in previous studies, including research on PI-RADS [15,21–23,27,28]. Third, even if MRI findings were negative, we performed a biopsy based on an elevated serum prostate-specific antigen level or abnormal digital rectal examination, which might have caused some selection bias. Fourth, slice thickness of all sequences should be ≤3 mm (the current study employed a 3.4-mm slice thickness). Fifth, this study was conducted to compare relaxometry measurements from synthetic MRI and the DCE-MRI PI-RADS score for csPCa diagnosis in category 3 lesions according to the literature of PI-RADS v2.1 guideline, while a comparison between these two tests in all lesions (i.e., not only category 3 lesions) was not included in this study. However, for both expert and basic prostate radiologists, the specificity of relaxometry measurements tended to be slightly lower than that of DCE-MRI in the evaluation of category 3 lesions. This trend may also be present in other lesions than only those in category 3. Future studies with larger sample sizes should assess if the diagnostic performance is comparable between relaxometry measurements and the DCE-MRI PI-RADS score in the evaluation of all lesions. Lastly, this prospective single-centre study comprised a relatively small sample size. Further larger, multicentre validation studies are necessary to confirm the results of this study.

6. Conclusions

In conclusion, the diagnostic performance of synthetic bpMRI was comparable to that of conventional bpMRI for primary PCa evaluation. Moreover, the diagnostic performance of relaxometry maps from synthetic MRI was similar to that of DCE-MRI PI-RADS score for csPCa in PI-RADS category 3 lesions in PZ, for both expert and basic prostate radiologists. Therefore, bpMRI with relaxometry maps derived from synthetic MRI shows potential as a contrast agent-free method for primary PCa evaluation.

Ethical Approval

Institutional Review Board approval was obtained.

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CRediT authorship contribution statement

Yuki Arita: Investigation, Writing – original draft; Hirotaka Akita: Resources, Data curation; Hirokazu Fujiwara: Resources, Data curation; Masahiro Hashimoto: Resources, Data curation; Keisuke Shigeta: Resources, Data curation; Thomas C. Kwee: Writing – review & editing; Soichiro Yoshida: Writing – review & editing; Takeo Kosaka: Resources, Data curation; Shigeo Okuda: Resources, Data curation; Mototsugu Oya: Supervision, Project administration; Masahiro Jinkaki: Supervision, Project administration.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.ejro.2022.100403.
