Comparison of optimised endovaginal vs external array coil T2-weighted and diffusion-weighted imaging techniques for detecting suspected early stage (IA/IB1) uterine cervical cancer

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Abstract

Objective To compare sensitivity and specificity of endovaginal versus external-array coil T2-W and T2-W+DWI for detecting and staging small cervical tumours.

Methods Optimised endovaginal and external array coil MRI at 3.0-T was done prospectively in 48 consecutive patients with stage Ia/Ib1 cervical cancer. Sensitivity/specificity for detecting tumour and parametrial extension against histopathology for a reading radiologist were determined on coronal T2-W and T2W+DWI images. An independent radiologist also scored T2-W images without and with addition of DWI for the external-array and endovaginal coils on separate occasions >2 weeks apart. Cohen’s kappa assessed inter- and intra-observer agreement.

Results Median tumour volume in 19/38 cases positive on subsequent histology was 1.75 cm³. Sensitivity, specificity, PPV, NPV were: reading radiologist 91.3 %, 89.5 %, 91.3 %, 89.5 %, respectively; independent radiologist T2-W 82.6 %, 73.7 %, 79.1 %, 77.8 % for endovaginal, 73.9 %, 89.5 %, 73.9 % for external-array coil. Adding DWI improved sensitivity and specificity of endovaginal imaging (78.2 %, 89.5 %); adding DWI to external-array imaging improved specificity (94.7 %) but reduced sensitivity (66.7 %). Inter- and intra-observer agreement on T2-W+DWI was good (kappa=0.67 and 0.62, respectively).

Conclusion Endovaginal coil T2-W MRI is more sensitive than external-array coil for detecting tumours <2 cm³; adding DWI improves specificity of endovaginal imaging but reduces sensitivity of external-array imaging.

Key Points

• Endovaginal more accurate than external-array T2-W MRI for detecting small cervical cancers.
• Addition of DWI improves sensitivity and specificity of endovaginal T2-W imaging.
• Addition of DWI substantially reduces sensitivity of external-array T2-W imaging.

Keywords Cervical cancer · Endovaginal coil · External array coil · Diffusion-weighted MRI · Fertility

Abbreviations

ADC Apparent diffusion coefficient
DWI Diffusion-weighted imaging
EPI Echo-planar imaging
FIGO Federation Internationale Gynecology and Obstetrics
FSE Fast spin-echo
LLETZ Large loop excision of the transformation zone
MRI Magnetic resonance imaging
SENSE Sensory encoding
SNR Signal-to-noise ratio
TSE Turbo spin-echo
Introduction

Increasingly, because of effective cytological screening programmes, uterine cervical cancer presents at an early stage with small-volume disease. A ‘see-and-treat’ policy at most centres results in referrals for imaging following a diagnostic cone biopsy or large loop excision of the transformation zone (LLETZ) procedure, at which a significant proportion of the disease may be excised. The presence of residual disease is critical to inform further surgical management especially if considering preservation of fertility, so that its identification has become a key requirement from imaging. Cervical amputation (trachelectomy) is an option in these cases, where fertility is not compromised for other reasons [1] but requires at least a 1-cm length of normal cervical canal between the superior tumour margin and the internal os [2, 3]. Therefore, knowledge of residual tumour margins in relation to the normal cervical anatomy [4, 5] is paramount when considering patients for trachelectomy. The superior soft-tissue contrast provided by magnetic resonance imaging (MRI) makes it the imaging technique of choice for demonstrating cervical tumour in relation to normal cervical anatomy [6] in preoperative imaging [7]; standard image acquisition using external phased array receiver coils at field strengths of 1.5-T or 3.0-T results in an in-plane resolution of >1 mm with a 3- to 4-mm slice thickness. At this resolution, tumours of >1 cm have a sensitivity and sensitivity of detection around 60 %, although specificity is high [8, 9], but smaller tumours can be missed because spatial resolution is limited by the signal-to-noise ratio (SNR) available.

The spatial resolution of MR images may be improved by the use of receiver coils close to the region of interest because they provide significant increases in SNR [10–12]. The use of an endovaginal receiver coil has previously been shown to improve the detection of tumours of <1.0 cm³ on T2-W imaging [13] and also allows detailed visualisation of the parametrium up to 6 cm from the surface of the internal coil [14]. When combined with diffusion-weighted (DW) contrast, the endovaginal technique further improves sensitivity and specificity of local staging to 92 % and 82 %, respectively [15, 16] because the apparent diffusion coefficient (ADC) of tumour is significantly lower than normal tissue [17]. However, a formal comparison of an optimised T2+DW protocol for tumour detection using an endovaginal versus an external array-only technique has hitherto not been undertaken because of the time and resource implications of doing both fully optimised protocols in the same patients prior to surgery.

The purpose of this work therefore was to undertake such a prospective study and compare the sensitivity and specificity of T2-W and T2-W+DWI obtained using an endovaginal coil with that using an external array coil for detecting small-volume cervical tumours and identifying early parametrial extension.

Methods

Endovaginal coil

A 37-mm ring-design solenoidal receiver coil similar to one previously described for use at 1.5-T [18] was used as part of a coil array with an externally placed flex-L coil (Philips Medical Systems, Best, The Netherlands). The endovaginal coil was designed and built in-house. It was covered in a disposable sheath and required simple manual insertion, without visual or imaging guidance, and manipulated so that the ring encompassed the cervix. The coil was immobilised in an external clamp as previously described. Vaginal air introduced during insertion was aspirated using a Ryles tube to minimise image distortion.

External array

A receive-only SENSE XL Torso coil 16-element phased array was used (Philips Medical Systems). This is a flexible volume coil consisting of anterior and posterior sections, each of eight elements – four upper and four lower. Coil dimensions were 74 mm × 558 mm × 600 mm for each half (upper and lower). Each element had a Z coverage of 22.5 cm.

Sequence optimisation

We initially established the most appropriate imaging protocols for both the endovaginal technique and the external array technique. As high a resolution as possible with the T2-W sequences without compromising image quality was deemed to be a minimum requirement. We matched the voxel size of the external array imaging to the endovaginal DW imaging (Table 1). From our experience in reporting these images, planes sagittal and coronal to the cervix were selected. The endovaginal images had a spatial resolution of 0.35 mm³ (0.42 × 0.42 × 2 mm), while the torso array had a spatial resolution of 3 mm³ acquired (1 × 1 × 3 mm). In order to minimise acquisition time, DWI for comparison was done in the coronal plane only as parametrial extension is best demonstrated in this plane. For the endovaginal technique we used the ZOOM sequence [19] which substantially reduces image distortions that arise due to susceptibility artefacts from the endovaginal coil. This allows a spatial resolution on the DW images of 3.1 mm³ (1.25 × 1.25 × 2 mm). Because of the lower image

| ZOOM | Zonal oblique multislice |
| PPV | Positive predictive value |
| NPV | Negative predictive value |
| RVT | Radical vaginal tracheectomy |
SNR without the internal coil, it was not possible to derive any meaningful data with the ZOOM-DWI. A conventional echo-planar imaging (EPI) sequence with a pixel resolution equivalent to that of the ZOOM-DWI image was therefore selected (1.25×1.25 mm) but a slice thickness of 3 mm was selected to match the slice thickness of the T2-W images and retain SNR, giving a voxel size of 4.7 mm³.

Details of the endovaginal coil and external array coil protocols are given in Table 1. SNR measurements were made in a test-object with each coil combination using the optimised sequences and corresponding reconstruction filters, multichannel reconstruction and parallel imaging. A 2.5-cm diameter gel test object (Eurospin, T1=500 ms, T2=50 ms) surrounded by mineral oil test objects to mimic a body load was imaged with the endovaginal-flex-L coil combination followed by the external array coil, repeating each sequence three times. Mean signal intensity measured from a 200 mm² region of interest on the central slice within the gel test object from the three repetitions was divided by the mean of the standard deviation of background noise. There was near equivalence of the SNR on T2-W images with each coil combination. There was a 62% reduction in SNR on the b=800 s/mm² ZOOM-DWI images from the endovaginal coil compared to the corresponding T2-W images, but a seven times reduction in SNR on the b=800 s/mm² DWI from the external array coil compared to the endovaginal DWI images (Table 1).

Patients

The study was approved by the local Institutional Review Board. All patients gave informed, written consent. Over a 15-month period (May 2013–Aug 2014), a further follow-on cohort from our previous study [15, 20] of 51 patients were referred for MRI following an abnormal smear test and subsequent histological evidence of invasive cervical cancer. Twenty-two complained of at least one episode of abnormal bleeding unrelated to any preceding surgical procedure. All were successfully prospectively recruited for assessment with endovaginal T2-W MRI followed by high-resolution external phased array MRI at 3.0-T. Three patients were staged as Federation Internationale Gynecology and Obstetrics FIGO 2B at examination under anaesthesia, so that 48 consecutive patients with Stage 1 cervical cancer aged 24–53 years (mean 34.4±8.2 years) were included in this study (Fig. 1). Forty-two patients had a cone or LLETZ biopsy at a median of 4.0 weeks (range 2–12 weeks) prior to MRI in which some or the entire tumour may have been excised; at histology, 26 were

### Table 1 Sequence parameters used for imaging

<table>
<thead>
<tr>
<th>Parameter</th>
<th>T2-weighted coronal, sagittal</th>
<th>DWI coronal</th>
<th>T2-weighted DWI</th>
<th>DWI</th>
</tr>
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<tbody>
<tr>
<td>Coil selection</td>
<td>Dual: Endovaginal+FLEX-L 2 element phased array body</td>
<td>Single: Endovaginal</td>
<td>SENSE-XL torso</td>
<td>SENSE-XL torso</td>
</tr>
<tr>
<td>Sequence</td>
<td>Turbo Spin Echo-Multi Shot (TSE factor 10)</td>
<td>Spin-Echo Echo Planar Imaging-Single Shot</td>
<td>TSE (TSE Factor 13)</td>
<td>EPI</td>
</tr>
<tr>
<td>TR (ms)</td>
<td>2750 (coronal) 2500 (sagittal)</td>
<td>6500 2100 3230</td>
<td>3.125 [1.25×1.25×2] 3.00 [1×1×3] 4.69 [1.25×1.25×3]</td>
<td></td>
</tr>
<tr>
<td>TE (ms)</td>
<td>80 54 90 90 90</td>
<td>217.7 217.3 9.6</td>
<td>2100 120 110×100</td>
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<tr>
<td>Flip angle</td>
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<td>90°</td>
<td>90°</td>
<td>90°</td>
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<tr>
<td>Bandwidth (Hz/Px)</td>
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<td>100×100</td>
<td>120×200 110×100</td>
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<td>FOV (mm)</td>
<td>4.7×4.7 4.7×4.7 4.7×4.7 4.7×4.7 4.7×4.7</td>
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<td>4.7×4.7 4.7×4.7</td>
<td></td>
</tr>
<tr>
<td>Acq pixel size (mm)</td>
<td>0.42×0.42 1.25×1.25 1.25×1.25</td>
<td>1.25×1.25</td>
<td>1.25×1.25 1.25×1.25</td>
<td></td>
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<td>Acq matrix</td>
<td>238</td>
<td>80 120 88</td>
<td>80 120 88</td>
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</tr>
<tr>
<td>Recon matrix</td>
<td>288</td>
<td>224 400 224</td>
<td>224 400 224</td>
<td></td>
</tr>
<tr>
<td>Acq voxel size (mm³)</td>
<td>0.35 [0.42×0.42×2] 3.125 [1.25×1.25×2] 3.00 [1×1×3] 4.69 [1.25×1.25×3]</td>
<td>0.35 [0.42×0.42×2] 3.125 [1.25×1.25×2] 3.00 [1×1×3] 4.69 [1.25×1.25×3]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Slice thickness (mm)</td>
<td>2 (0.1 separation) 2 (0.1 separation)</td>
<td>2 (0.1 separation)</td>
<td>2 (0.1 separation) 2 (0.1 separation)</td>
<td></td>
</tr>
<tr>
<td>Recon voxel size (mm³)</td>
<td>0.24 [0.35×0.35×2] 0.40 [0.45×0.45×2] 0.75 [0.5×0.5×3] 0.72 [0.49×0.49×3]</td>
<td>0.24 [0.35×0.35×2] 0.40 [0.45×0.45×2] 0.75 [0.5×0.5×3] 0.72 [0.49×0.49×3]</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of slices</td>
<td>24 (coronal) 24 (sagittal)</td>
<td>24</td>
<td>24 16 16</td>
<td></td>
</tr>
<tr>
<td>NSA</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Included b-values s/mm²</td>
<td>n/a 0, 100, 300, 500, 800 n/a 0, 100, 300, 500, 800</td>
<td>n/a 0, 100, 300, 500, 800</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Acquisition time</td>
<td>4 min 18 s (coronal) 4 min 0 s (sagittal)</td>
<td>4 min 33 s 3 min 58 s</td>
<td>4 min 18 s (coronal) 4 min 0 s (sagittal)</td>
<td></td>
</tr>
<tr>
<td>Signal in gel (T1=500 ms, T2=50 ms) to background noise</td>
<td>Flex-L elements 15 cm apart, gel sited centrally within endovaginal coil ring</td>
<td>Gel sited centrally within endovaginal coil ring</td>
<td>Coil elements 15 cm apart, gel sited centrally 51</td>
<td>Coil elements 15 cm apart, gel sited centrally 51</td>
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<tr>
<td></td>
<td>135 51</td>
<td>130 51</td>
<td>130 51</td>
<td></td>
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</tbody>
</table>

**DWI** diffusion-weighted imaging, **NSA** number of signal averages
squamous cell carcinoma and 16 adenocarcinoma. The other five had punch biopsies only (three squamous cell carcinoma, two adenocarcinomas); of these three had undergone cone or LLETZ biopsies more than 9 months previously.

**Imaging methods**

Images were obtained at 3.0-T (Philips Achieva). Hyoscine butyl bromide (Buscopan) 20 mg intramuscularly (IM) was administered to reduce artefact from bowel peristalsis. Endovaginal imaging was performed first. T2-W images were acquired in three orthogonal planes to the cervix and covered it entirely. DWI slice-matched images were acquired in free-breathing coronal to the cervix as parametrial extension is best assessed in this plane. Sequence parameters and acquisition time are given in Table 1. Isotropic apparent diffusion coefficient (ADC) maps based on the monoexponential rate of signal decay with increasing diffusion weighting were generated with the system software (Philips Medical Systems) using all b-values.

Following the endovaginal imaging, the coil was removed and, following a 5-minute off-bed break, external array coil imaging was done. T2-W images were acquired in planes sagittal and coronal to the cervix. DW-slice-matched images were obtained coronal to the cervix. As with the endovaginally acquired data, isotropic ADC maps based on the monoexponential rate of signal decay with increasing diffusion weighting were generated with the system software (Philips Medical Systems) using all b-values.

**Data analysis**

At the time of clinical image reading, the reporting radiologist (20 years’ experience of pelvic MRI), with access to clinical information and all imaging data sets noted the absence or presence of tumour and the absence or presence of tumour invasion into the right and left parametrium. Where tumour was noted, a tumour volume was measured by drawing a region-of-interest around the tumour on every slice on which it appeared and multiplying the total measured tumour area by the slice thickness plus interslice gap.

On completion of the patient accrual, an independent radiologist (5 years’ experience of pelvic MRI) without access to the clinical data separately scored first the anonymised coronal and sagittal T2-W external array images as positive or negative for tumour within the cervix; the absence or presence of parametrial extension was also noted as absent or present. This was then repeated with inclusion of the corresponding coronal ADC maps. At a subsequent occasion 2 weeks later to avoid memorisation, the same observer scored the randomized, anonymised coronal and sagittal T2-W endovaginal images without and then with the addition of the corresponding coronal ADC maps. The absence or presence of right and left parametrial extension was also noted.

**Histopathological analysis**

Tissue was acquired at radical hysterectomy (n=10), radical vaginal trachelectomy (RVT) (n=13), cold knife cone (n=11) or LLETZ procedure (n=4) in 38 patients. Of the

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**Fig. 1** Flow chart of patients included in the study and their clinical pathway. MRI magnetic resonance imaging, DW-MRI diffusion-weighted MRI, FIGO Federation Internationale Gynecology and Obstetrics, LLETZ Large loop excision of the transformation zone
remainder, four went to chemoradiotherapy and were known to have prior positive biopsies and six were deemed to have negative MRI scans with clear margins on the initial cone or LLETZ biopsy so no further intervention at that point was deemed necessary. These latter six cases were therefore excluded from the analysis because of a lack of post-MRI confirmation of the absence of tumour. All specimens were fixed in formalin. For hysterectomy and RVT specimens, a shave of the proximal endocervical resection margin (at or near the internal os) was sliced longitudinally (sagittal plane centrally, coronal plane laterally). The cervix together with the parametrium was sliced into transverse sections up to approximately 1 cm from the ectocervix. The remaining 1 cm of distal cervix together with vaginal cuff was sectioned longitudinally (sagittal plane centrally, coronal plane laterally or sagittally throughout). All slices (each 3–4 mm thick) were processed and embedded in paraffin; 2–3 micron sections were deparaffinised and stained with haematoxylin and eosin. According to the thickness of the slice obtained at macroscopy, each histology slide was separated from the next by 3–4 mm.

Statistical analysis

Sensitivity, specificity, and positive and negative predictive values for detecting the presence of tumour by the reading radiologist were obtained. For the independent radiologist blinded to the clinical information sensitivity, specificity, and positive and negative predictive values of T2-W without and with the addition of DWI were established for detecting the presence of tumour. Data were obtained for the endovaginal technique and separately for the external array-only technique. Similar data were derived for right and left parametrial extension. A Cohen’s kappa to assess agreement between observers for interpreting endovaginal T2-W + DWI imaging together was calculated between reading and independent radiologists and between the internal and external coil readings of the independent radiologist on T2-W and separately T2-W + DWI imaging.

Results

Thirty-eight patients (ten Stage Ia, 28 Stage 1b) underwent a surgical procedure a median of 27 days (lower quartile 23 days, upper quartile 35 days) after MRI; of these, 19 had residual tumour and 19 had no residual tumour on histology. Four patients who received chemoradiotherapy had tumour on biopsy at staging examination under anaesthesia (Fig. 1). Median tumour volume measured on endovaginal MRI of 23 true positive lesions (Fig. 2) was 1.75 cm³ (lower quartile 0.77 cm³, upper quartile 5.5 cm³).

Sensitivity and specificity for detecting tumour by the reporting radiologist were 91.3 % and 89.5 %, respectively. There were two false positives (Fig. 3) with tumour volumes estimated as 0.05 cm³ and 1.5 cm³, the latter described as a flat ectocervical plaque. The two false negative lesions (Fig. 4) were 6 mm in maximum dimension on histology. Sensitivity and specificity for detecting tumour by the independent radiologist blinded to the clinical information on endovaginal T2-W alone was 82.6 % and 73.7 %; inclusion of the DWI images showed a small reduction in sensitivity to 78.2 % but improved specificity to 89.5 %. In comparison, the external array T2-W images showed a lower sensitivity for detecting tumour of 73.9 % but a high specificity of 89.5 %. Addition of the DWI images improved specificity further to 94.7 % but at the expense of a substantial loss in sensitivity to 66.7 % (Table 2). The agreement between observers for interpreting endovaginal T2-W + DWI images together was good (Cohen’s kappa = 0.67). Agreement between the independent radiologist assessment of the T2-W images on internal and external techniques was moderate (kappa = 0.49), while the assessment of the T2-W + DW images on internal and external techniques was good (kappa = 0.62).

In 26 cases who went for trachelectomy or radical hysterectomy, one case was at a different institution and information on the parametrial status was not available. Therefore, 50 parametrial regions were evaluated.
in 25 cases, assuming that right and left parametrial extension are effectively independent. Tumour within the cervix had been noted on MRI in 17 of these cases by the reporting radiologist and in 17 by the independent radiologist. Tumour within a normal-sized parametrial node was present in one case only on histology. This was missed by the reporting radiologist who marked this parametrial region as negative. The sensitivity, specificity, PPV and NPV for the independent radiologist using the endovaginal compared to the external array technique, and without and with the DWI, are given in Table 3. However, as we had only one case with parametrial extension in our cohort, no firm conclusions can be drawn about the differences in parametrical assessment with and without an endovaginal coil.

Discussion

These data clearly demonstrate a higher sensitivity of T2-W endovaginal imaging for detecting small-volume tumours compared to an external array technique, albeit with lower specificity. Importantly, however, the addition of DWI to the endovaginal imaging serves to improve specificity so that it is equivalent to that of the external array T2-W assessment, but the addition of DWI to the external array technique substantially reduces sensitivity of detection of these small lesions because the spatial resolution of DWI with external array imaging is limited by low SNR. The higher sensitivity of detection of small tumours on endovaginal T2-W imaging alone compared to external array coil imaging is to be expected because of the improved spatial resolution with the increased SNR available with a small coil in situ, and the correlation of
tumour size with histology has been demonstrated [20]. This has previously been shown not just in endovaginal imaging [18, 21] but also in prostatic imaging [22] where an endorectal technique is now standard in clinical practice.

The improved SNR afforded by an endovaginal technique also means that sequences inherently low in SNR can be more usefully deployed. In prostatic imaging the equivalent endorectal imaging has led to the routine acquisition of

**Fig. 4** False negative on endovaginal T2-W with negative diffusion-weighted imaging (DWI) in a 27-year-old female: Sagittal (a), axial (b), coronal (c) T2-W (FSE TR 2750 ms, TE 80 ms) and ADC map (EPI, TR 6500 ms, TE 54 ms, b=0, 100, 300, 500, 800 mm²/s, d) with corresponding external array coil T2-W coronal (FSE TR 2100 ms, TE 90 ms, e) and ADC map (EPI TR 3230 ms, TE 53 ms, b=0, 100, 300, 500, 800 mm²/s, f). There was no evidence of tumour on MRI with either technique, but a 6-mm microinvasive tumour (Stage Ia) was noted on histopathology (g, arrow)
Another approach to improving geometric distortions on the DW images [27] is to replace the echo-planar based sequences with a turbo spin-echo (TSE). Previous attempts at such an approach have resulted in successful implementation of turbo spin-echo DWI for head and neck [28] and endometrial [29] cancers. The resulting images are more robust and suffer less from distortion, but incur a time penalty. Typically an increase of around 50% in imaging time is expected. There is also an increased tendency to motion artefact and as these images are not single-shot like their echo-planar based counterpart, there is increased image blurring [30]. Optimisation of the DW technique for endovaginal MRI is therefore warranted; evaluation of the advantages and limitations of an EPI versus TSE-based sequences is required but was beyond the scope of the study, not least because of the length of a detailed endovaginal examination followed by a detailed external array examination. The number of b-values to use is also debated [31], but the choice of five b-values in this instance was based

**Table 2** Sensitivity, specificity, and positive (PPV) and negative (NPV) predictive values for detecting tumour by a reporting radiologist and an independent radiologist. The comparison of endovaginal vs. external array techniques without and with the addition of diffusion-weighted (DWI) magnetic resonance imaging to the T2-W sequences for the independent radiologist is shown

<table>
<thead>
<tr>
<th>N=42</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>PPV</th>
<th>NPV</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Reporting rad. T2W+DWI</td>
<td>91.3%</td>
<td>89.5%</td>
<td>91.3%</td>
<td>89.5%</td>
</tr>
<tr>
<td>T2-W</td>
<td>86.2%</td>
<td>73.7%</td>
<td>79.1%</td>
<td>77.8%</td>
</tr>
<tr>
<td>Independent rad. endovaginal</td>
<td>89.5%</td>
<td>73.9%</td>
<td></td>
<td></td>
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<tr>
<td>Independent rad. external array</td>
<td>89.5%</td>
<td>73.9%</td>
<td></td>
<td></td>
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</tbody>
</table>

**Table 3** Sensitivity, specificity, and positive (PPV) and negative (NPV) predictive values for detecting parametral extension by an independent radiologist. The comparison of endovaginal vs. external array catheter techniques with and without the addition of diffusion-weighted magnetic resonance imaging (DW-MRI) to the T2-W sequences is shown

<table>
<thead>
<tr>
<th>N=50 regions (25 patients)</th>
<th>Sens</th>
<th>Spec</th>
<th>PPV</th>
<th>NPV</th>
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<tr>
<td></td>
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<tr>
<td>T2-W</td>
<td>100%</td>
<td>100%</td>
<td>100%</td>
<td>100%</td>
</tr>
<tr>
<td>Independent rad. endovaginal</td>
<td>100%</td>
<td>95%</td>
<td>33.3%</td>
<td>100%</td>
</tr>
<tr>
<td>Independent rad. external array</td>
<td>100%</td>
<td>97.9%</td>
<td>50</td>
<td>100%</td>
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References


Conclusion

Sensitivity for detecting tumour within the cervix was higher with an endovaginal coil technique than with a conventional external array coil, although specificity was lower. The addition of DWI to the endovaginal T2-W imaging gave a small reduction in sensitivity, but with a substantial improvement in specificity, making this the preferred diagnostic option. Although less sensitive, specificity for tumour detection was high with the external array coil using T2-W imaging; the addition of DWI in this case did not improve specificity, but markedly reduced sensitivity. Therefore DWI is only useful in early stage cervical cancer as an adjunct to T2-W imaging if an endovaginal technique is used. Endovaginal coil MRI would be recommended for assessment of clinical stage 1a and 1b1 cervical cancer.

on the ability to calculate ADC more accurately, even though quantitative assessments did not form part of the current work.

The scan schedule in this small group of patients was intensive and took ~60 min but fortunately would not be required in standard clinical care. It is our normal practise to do a detailed evaluation of the cervix using the endovaginal coil over 25 min and to then remove it and assess the pelvic and abdominal lymph node chains with an external array, which takes another 20 min. Although longer than a standard external array pelvic MRI of 30 min, this has the added value of detailed data prior to planning a fertility-sparing surgical procedure such as a trachelectomy in these young women.

Appropriate referral for an endovaginal MRI is largely based on an approximate clinical or histological assessment of tumour volume. Larger tumours, particularly stage 2b and above, are adequately imaged using an external array coil technique [32, 33]. Tumours benefiting from this procedure are predominantly those clinically staged as Ia2 or Ib1, where the tumour may or may not be just palpable or the diagnostic or therapeutic cone biopsy or LLETZ indicates that there is incomplete excision of tumour or tumour extending to margins of the specimen. In these cases, endovaginal MRI is invaluable in depicting the presence of residual tumour and defining its extent in relation to the endocervical canal pre-operatively. As referrals to our study were largely based on these criteria, the tumour median volume in our cohort was 1.75 cm³. The increased detectability of these small tumours with endovaginal MRI is directly related to the improved spatial resolution possible because of the substantial improvement in SNR afforded by the internal coil. Although the numbers of patients are small, the sensitivity and specificity of our endovaginal reads by the reading radiologist are identical to those of previous cohorts studied with this technique [20].