Magnetic Resonance Imaging of Uterine Cervix: A Pictorial Essay

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Published online: 2021-08-04

Abstract

Uterine cervix is the lower constricted part of uterus which is best evaluated by magnetic resonance imaging (MRI) due to its higher soft tissue and contrast resolution. The cervical cancer is a common gynecological cancer causing much morbidity and mortality especially in developing countries. Cervical carcinomas mainly occurs in reproductive age group with prognosis mainly depending on the extent of disease at the time of diagnosis, hence it is important to identify these cancerous lesions early and stage them accurately for optimal treatment. In this article, we will review the following: (1) the normal MRI anatomy of uterine cervix; (2) MRI protocol and techniques in evaluation of cervical lesions; (3) imaging of spectrum of various congenital abnormalities and pathologies affecting uterine cervix which ranges from congenital abnormalities to various benign lesions of cervix like nabothian cysts, tunnel cysts, cervicitis, cervical fibroid, and, lastly, endometriosis which usually coexists with adenomyosis; the malignant lesions include carcinoma cervix, adenoma malignum or direct extension from carcinoma endometrium or from carcinoma of vagina; (4) Accurately stage carcinoma of cervix using FIGO classification (2018); and (5) posttreatment evaluation of cervical cancers. MRI is the most reliable imaging modality in evaluation of various cervical lesions, identification of cervical tumors, staging of the cervical malignancy, and stratifying patients for surgery and radiation therapy. It also plays an important role in detection of local disease recurrence.

Keywords

► cervical lesions
► cervical cancer
► MRI of cervix

Introduction

Uterine cervix is the lower constricted part of uterus which is best evaluated by magnetic resonance imaging (MRI) due to its higher soft tissue and contrast resolution. In this article, we shall evaluate the normal anatomy of cervix; optimal MRI techniques; and role of various sequences, congenital anomalies of cervix, benign, and malignant lesions involving cervix.

Anatomy

The word “cervix” means neck in Latin, is the lower fibromuscular portion of the uterus, cylindrical or conical in shape. The cervix projects into the vagina through the upper part of its anterior wall and opens through external opening, that is, external os. The upper half of cervix, above the vagina, is called supravaginal portion which is attached to the surrounding tissues, whereas the lower free portion...
of cervix called portio vaginalis is limited by anterior and posterior vaginal fornices. The portio is further divided into anterior and posterior lips of which anterior lip is shorter and projects lower than posterior lip. The internal os (isthmus) is located between the supravaginal portion of cervix and uterine corpus. The ectocervix is the portion of cervix lying external to external os and is visualized on per speculum examination. The endocervix is proximal to external os and endocervical canal extends from internal to external os connecting the uterine cavity with the vagina. Maximum width of endocervical canal is 8 mm. The endocervix is lined with columnar epithelium as compared with ectocervix which has nonkeratinizing squamous epithelium. With age, the squamous cells cover the columnar cells of endocervical glands resulting in superior migration of the squamocolumnar junction. This transitional area is most prone to development of carcinoma.

The cervix measures approximately 3 to 4 cm in length, and 2.5 cm in diameter; however, its size and shape vary depending on the woman’s age, parity, and hormonal status. Before the onset of menstruation, the cervix is about half to two-thirds of that of uterus. In adult females, cervix constitutes about one-third of uterus, while in postmenopausal females, its length is almost same as uterine corpus.\(^1,2\)

**Vascular Supply**

Blood supply to cervix is by descending branches of uterine arteries which are branches of anterior division of internal iliac arteries. These arteries reach the lateral cervical walls along the upper margins of cardinal ligaments. The venous drainage is parallel to arterial supply with communication with cervical plexus and neck of urinary bladder. The lymphatic drainage is to external iliac, obturator, hypogastric, common iliac, sacral nodes, and nodes of the posterior wall of urinary bladder.\(^3\)

**Magnetic Resonance Imaging Technique and Imaging Protocol**

**Patient Preparation**

The scan is performed with the patient fasting for at least 4 hours to reduce bowel peristalsis artifacts. Patient is asked to empty the bladder and bowel before examination to reduce blurring from motion artifacts and ghosting due to bowel peristalsis and bladder motion.\(^4\) Intramuscular or intravenous administration of the antiperistaltic agents appears to improve image quality and lesion visualization in oncological pelvic MRI; however, it is not a routine practice at our center. Sterilized gel is used to distend the vagina for better evaluation of cervical lesions. MRI should be performed before biopsy or at least 10 days after biopsy to avoid false positive results due to local inflammation.\(^5\)

**Magnetic Resonance Imaging Protocol**

A phase-array surface coil is used to increase signal-to-noise ratio (SNR). We use following sequences on 3T Philips Achieva MRI Scanner:

1. Axial T2-weighted (T2W) fat saturated with large field of vision (FOV).
2. Axial T1-weighted (T1W) with large FOV.
3. Thin sagittal T2W.
4. Thin coronal T2W.
5. Thin axial T2W.
6. Axial T1 fat saturated (if needed to differentiate fat from hemorrhage).
7. Diffusion-weighted images (DWIs).
8. Susceptibility-weighted images (SWIs; if required).

We first obtain T2 fat-saturated and T1W Turbo Spin Echo sequence in the true axial plane of the pelvis with large FOV to have global view of pelvis (►Fig. 1). High-resolution, small FOV, 3- to 4-mm thick with zero-gap sections of T2Weighted sequences are obtained in the sagittal, oblique axial, and oblique coronal planes which are perpendicular and parallel to cervix (►Fig. 2). Sagittal T2W sequence is useful in assessing the location of lesion in cervix (anterior or posterior), its relation with uterine body, vagina, as well as with rectum and urinary bladder. Involvement of internal os and distance of tumor/lesion from internal os is also best seen on this plane.

Oblique axial T2W sequence is useful in assessment of parametrial and lateral pelvic invasion and bowel, bladder, and ureteric involvement with proximal hydroureter. Pelvic lymph nodes are well seen on true axial and coronal T2W images, as well as on DWIs; however, they are best visualized on T2 fat-saturated axial images with large FOV.

Oblique coronal plane assesses the invasion of parametrium well.\(^7\)

T1W images provide information on pelvic anatomy, help in evaluation of lymph nodes, and bone marrow. Hemorrhage in lesion appears hyperintense on T1W images and T1 fat-saturated images are helpful to differentiate it from fat.

We also perform DWI and SWI routinely in all the cases. DWI is extremely helpful in assessment of cellularity of the tumors. Cervical carcinomas show restricted diffusion, that is, hyperintense signal on DWI and hypointense signal on ADC maps, thus facilitating the exact extent of the tumor.
DWI is also useful in assessing parametrial/pelvic invasion, nodal metastases, and recurrent/residual disease. Keriakos and Darwish used cut-off ADC value of $1.12 \times 10^{-3}$ with specificity of 100% and sensitivity of 87.5% to distinguish benign and cervical lesions.

Routine MRI with DWI in combination can accurately distinguish between benign and malignant cervical lesions and is preferable to dynamic contrast MRI. Use of contrast is not necessary in staging of cervical cancer, as noncontrast images provide all the necessary information. Dynamic contrast-enhanced T1 fat-saturated images are mainly performed for endometrial carcinoma and are not part of routine protocol for staging of cervical carcinoma; however, these may be considered in evaluation of small cervical tumors, lesions with equivocal T2 findings, and in distinguishing tumor recurrence from posttreatment changes.

**Normal Magnetic Resonance Imaging Anatomy**

Cervix on T2W images shows the following distinct zones from within outside (Fig. 2):

- Zone 1: central hyperintense mucous in cervical canal.
- Zone 2: high-signal intensity endocervical mucosa and glands (contains numerous folds and clefts as the pli-cae palmate). Combined thickness of zones 1 and 2 is 2 to 3 mm.
- Zone 3: hypointense fibrous stroma is 3- to 8-mm thick.
- Zone 4: outer intermediate signal intensity of loose stroma.

On T1W images, cervix appears as intermediate signal intensity structure without clear visualization of its zonal anatomy. The uterine corpus is smaller than the cervix in premenstrual girls (Fig. 3), whereas after menopause, uterus as a whole is smaller and atrophied with cervical length almost equal to uterine body (Fig. 4). Postoperative low-segment caesarean section (LSCS) scar is seen a thin-linear T2 hyperintense signal with associated muscular loss and thinning of anterior uterine wall (Fig. 5).

**Pathologies of Cervix**

**Congenital Anomalies**

Uterus, cervix, fallopian tubes, and upper two-thirds of vagina arise from paired Müllerian ducts which eventually undergo fusion and resorption in utero. Any disruption in this process results in a spectrum of congenital anomalies termed as Müllerian duct anomalies (MDA). MRI is commonly used for clinical practice for confirmation when these anomalies are suspected on ultrasonography or hysterosalpingography.

The most widely accepted classification of MDAs is of the American Society for Reproductive Medicine system (Table 1). This classification is a framework and not all anomalies completely fit into its seven categories.
Embryogenesis: female genital tract development occurs in three main stages, that is, ductal development, ductal fusion, and ductal degeneration. Hence, the anomalies are also characterized due to nondevelopment, nonfusion, and nondegeneration. We divided the cervical anomalies as seen on MRI into three groups: aplasia or hypoplasia of cervix, cervical duplication, and septum in cervix (Table 2).

1. Agenesis or hypoplasia: these anomalies occur when there is early failure of ductal development. Mayer–Rokitansky–Kuster–Hauser syndrome is characterized by agenesis of proximal vagina, cervix, and uterus with normal ovaries very well seen on MRI (Figs. 6 and 7). They present with primary amenorrhea. Few cases of Turner’s syndrome (partial or complete absence of X chromosome [45 XO]) can rarely have Müllerian agenesis with absent ovaries. Usual MRI findings in Turner’s syndrome include well-formed uterus, cervix, and vagina which are smaller in volume (hypoplastic) with streak ovaries (Fig. 8). Unicornuate uterus (class II) is characterized by normal development of one horn and partial or complete nondevelopment of other horn. The rudimentary horn, if present, may be without uterine cavity or in presence of cavity may be communicating or not communicating with the normal side. MRI shows a small curved unicornuate uterus and small cervix with normal zonal anatomy seen off midline. A noncommunicating rudimentary horn may show presence of hematometra and hematosalphinx secondary to endometriosis (Figs. 9 and 10).

2. Duplication of cervix: this can occur in uterus didelphys (class III) which is complete failure of fusion resulting in duplication of uterine horns, cervix, and proximal vagina and MRI shows duplicated two uteri, two cervices separated by normal myometrium (Fig. 11). Duplicated proximal vagina may be associated with transverse septum resulting in hematocolpos. Bicornuate uterus (class IV) which is partial failure of fusion and may be associated with duplicated cervix when muscular septum extends up to external os (bicornuate bicollis) or single cervix when septum is seen up to internal os (bicornuate unicollis).

3. Septum in cervix: this is seen in septate uterus (class VI) which is the most common congenital uterine anomaly occurring due to partial or complete nonreabsorption of the uterovaginal septum. The septum can be fibrous or muscular. Septate uterus is characterized by convex external fundal contour and apex of external fundal contour is more than 5-mm above interostial line to differentiate from fusion anomalies. Intercornual distance of less than 4.0 cm has been used to distinguish a septate from a bicornuate uterus.

Table 2 Cervical anomalies based on MRI appearance

<table>
<thead>
<tr>
<th>MRI appearance</th>
<th>MDA class</th>
</tr>
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<tbody>
<tr>
<td>Absent or hypoplastic</td>
<td>Includes classes I and II MDA</td>
</tr>
<tr>
<td>cervix</td>
<td></td>
</tr>
<tr>
<td>Duplication of cervix</td>
<td>Includes classes III and IV MDA</td>
</tr>
<tr>
<td>Septum in cervix</td>
<td>Class V MDA</td>
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</tbody>
</table>

Abbreviation: MDA, Müllerian duct anomalies; MRI, magnetic resonance imaging.

Table 1 Classification of MDAs by American Society for Reproductive Medicine system

<table>
<thead>
<tr>
<th>Class</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Segmental agenesis and variable degrees of uterovaginal hypoplasia</td>
</tr>
<tr>
<td>Ia:</td>
<td>vaginal</td>
</tr>
<tr>
<td>Ib:</td>
<td>cervical</td>
</tr>
<tr>
<td>Ic:</td>
<td>fundal</td>
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<tr>
<td>Id:</td>
<td>tubal</td>
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<tr>
<td>Ie:</td>
<td>combined</td>
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<tr>
<td>II</td>
<td>Unicornuate uterus which is characterized by partial or complete unilateral hypoplasia</td>
</tr>
<tr>
<td>IIa:</td>
<td>communicating rudimentary horn</td>
</tr>
<tr>
<td>IIb:</td>
<td>noncommunicating rudimentary horn</td>
</tr>
<tr>
<td>IIc:</td>
<td>no cavity</td>
</tr>
<tr>
<td>IId:</td>
<td>no horn</td>
</tr>
<tr>
<td>III</td>
<td>Uterus didelphys in which duplication of the uterus results from complete non-fusion of the Müllerian ducts.</td>
</tr>
<tr>
<td>IV</td>
<td>Bicornuate uterus which is characterized by incomplete fusion of the superior segments of the uterovaginal canal</td>
</tr>
<tr>
<td>IVa:</td>
<td>complete</td>
</tr>
<tr>
<td>IVb:</td>
<td>partial</td>
</tr>
<tr>
<td>V</td>
<td>Septate occurs due to nonresorption of uterovaginal septum</td>
</tr>
<tr>
<td>Va:</td>
<td>complete</td>
</tr>
<tr>
<td>Vb:</td>
<td>partial</td>
</tr>
<tr>
<td>VI</td>
<td>Arcuate results from near complete resorption of the septum</td>
</tr>
<tr>
<td>VII</td>
<td>DES associated uterine anomalies</td>
</tr>
</tbody>
</table>

Abbreviation: MDA, Müllerian duct anomalies.
Fig. 6 Congenital: uterine and cervical aplasia: sagittal (A) and axial (B) T2W images in 18-year-old girl with primary amenorrhea shows complete absence of uterus including cervix. Lower vagina is seen (arrow). Both ovaries appear normal with multiple follicles (asterisk). These findings are suggestive of Meyer–Rokitansy–Kuster–Hauser syndrome. T2W, T2-weighted.

Fig. 7 Congenital: cervical hypoplasia: sagittal (A), axial (B), and coronal (C) T2W images in a 14-year-old girl with primary amenorrhea shows complete absence of uterus. Small hypoplastic cervix and vagina are visualized (arrow). Both ovaries appear normal with multiple follicles (elbow arrows). T2W, T2-weighted.

Fig. 8 Turner’s syndrome: sagittal (A) and coronal (B and C) T2W images in a 26-year-old female with known Turner’s syndrome show well-formed uterus and cervix (asterisk); however, these are small in size. Both the ovaries are identified and also show smaller volumes (arrow). T2W, T2-weighted.

Fig. 9 Congenital: unicornuate uterus: sagittal (A and B) and axial (C) T2W images showing small unicornuate uterus with cervix on right side (arrow). Noncommunicating functional left rudimentary horn is seen showing blood-fluid level (asterisk). Ovaries are normal (elbow arrow). T2W, T2-weighted.

Fig. 10 Congenital: unicornuate uterus: coronal (A and B) T2W images showing small left unicornuate uterus (arrow). Few endometriotic cysts are seen in left ovary (asterisk). Rudimentary right horn (elbow arrow) is seen without endometrial cavity (nonfunctioning). T2W, T2-weighted.

Fig. 11 Congenital: uterus didelphys: coronal (A–C) T2W images in a 37-year-old female showing two widely divergent uterine horns (arrows) separated by a deep fundal cleft. Two separate cervices are present (elbow arrow). Two vaginas (asterisk) are also seen with vertical septum. T2W, T2-weighted.

Fig. 12 Benign lesions: nabothian cysts: sagittal (A), coronal (B) and axial (C) images showing multiple well-defined cysts in anterior as well as posterior lips of cervix (arrow).

Fig. 13 Benign lesions: cervicitis: coronal (A) and axial (B) T2W images in a 38-year-old female with Mirena intrauterine device for 4 years had symptoms of pain and whitish vaginal discharge. MRI shows Intra uterine contraceptive device (arrow) in situ and multiple tiny cystic lesions in cervix (elbow arrow) suggestive of cervicitis. MRI, magnetic resonance imaging; T2W, T2-weighted.
Benign Lesions

Nabothian cysts: these are retention cysts in cervix, formed as a result of healing of chronic cervicitis. In chronic cervicitis, there is proliferation of squamous epithelium which covers the columnar epithelium. Mucus secreted by columnar epithelium cannot be extruded out and results in formation of retention cyst. These are usually 2 to 10 mm in size and located on surface of endocervix; occasionally, they may be larger in size.7 Incidentally seen on MRI, these appear as well-defined small cystic lesions having intermediate or slightly high signal on T1W and very high on T2W images (Fig. 12). It may be a single-cyst or multiple-cystic lesion, located in endocervical lesion and occasionally extending into deep cervical stroma resulting in enlargement of cervix.9 Larger cysts can cause pressure symptoms due to compression of urinary bladder/rectum, pelvic pain, or can rupture resulting in vaginal discharge. It is necessary to mention number and size of cysts in report as these are benign and assure patient and physician. There can be entrapment of nabothian cysts post subtotal hysterectomy, resulting in formation of large cysts which can be symptomatic.

Tunnel cysts: tunnel cluster, a special type of nabothian cyst, is characterized by complex multicystic dilatation of the endocervical glands.7 These are benign, pseudoneoplastic cyst, is characterized by complex multicystic dilatation of cervical glands. These are seen in premenopausal women and are associated with chronic inflammation. Trichomonas vaginalis, Candida albicans, and herpes simplex virus cause inflammation of squamous epithelium of ectocervix, whereas Neisseria gonorrhoeae and Chlamydia trachomatis affect the columnar epithelium. Vaginal discharge with pelvic pain are the common presenting complaints.7 MRI is not indicated; however, if done, it may show some retention cysts in cervix which are non-specific (Fig. 13). It may also appear as multicystic lesion with hyperintense signal on T1W and T2W images due to presence of secretions/proteinaceous contents/blood products. Healing results in fibrosis often resulting in cervical stenosis.

Lobular endocervical glandular hyperplasia: this lesion usually occurs in women of reproductive age group; however, may occur in few postmenopausal women.19 It is an incidental finding, but can present as erosion or polyp with watery discharge being the presenting complaint. It is a benign hyperplastic lesion characterized by cystic dilatation of cervical glands.

On MRI, enlarged inner T2/DWI hypointense solid cervical stroma is observed along with central tiny cyst surrounded by multiple large cysts in periphery giving cosmos flower sign.20,21 The MRI appearance closely mimics adenoma malignum and it may be extremely difficult to differentiate these on MRI alone. Presence of mild glandular atypia and gastrin mucin along with MRI findings of multicystic lesion with inner solid component showed accuracy of 92% in a study by Takatsu et al.22 In comparison, solid lesion with atypical glandular cells was indicative of adenoma malignum or adenocarcinoma.

Cervical leiomyoma: these are extremely uncommon as compared with leiomyomas occurring in uterine body. Tiltman found cervical leiomyoma in 0.4% hysterectomies as compared with 64.6% in uterine myometrium.23 They may occasionally form polypoidal lesion and protrude into the cervical canal or vaginal canal and can cause maternal dystocia. They may also cause dysmenorrhea and hypermenorrhoea.7 On MRI, these appear as a well-defined, rounded, T2 hypo-/isointense lesion showing homogenous enhancement centered on cervix. Cellular leiomyomas have hyperintense signal on T2W images. However, various types of degeneration result in altered signal intensity (Figs. 14 and 15). Cervical leiomyomas may have unusual growth pattern, ill-defined border, and may enclose the cervical canal into the leiomyoma.23,24

Cervical polyps: cervical polyps are the most common cause of intermenstrual vaginal bleeding. Other presenting symptoms include menorrhagia, postcoital bleeding, postmenopausal bleeding, and vaginal discharge. Majority of these polyps are asymptomatic and incidentally detected on routine gynecological examination, or on ultrasound, or as a filling defect on hysterosalpingography. Hysteroscopy accurately determines the number, location and size of these polyps, and also can be therapeutic in hysteroscopic polypectomy.25

These are seen in premenopausal women and are associated with chronic inflammation. They may be pedunculated or sessile, well-circumscribed masses, with or without cysts filling the endocervical and vaginal canal.7 Histologically,
polyps encompass spectrum of various pathologies including endocervical/endometrial tissue with metaplasia. Other polypoidal cervical lesions include submucosal leiomyoma, malignancy, or blood clot. It has to differentiated from endometrial polyp by identifying the stalk with attached to the cervical wall; however, in 25% patients, both endometrial and cervical polyps may coexist. MRI accurately assesses cervical polyps or giant endometrial polyps extending into the cervical canal (►Fig. 16).

**Adenomyosis/endometriosis**: adenomyosis is characterized by ectopic endometrial tissue in myometrium, whereas endometriosis is presence of functional endometrial glands and stroma outside the uterine cavity and myometrium. The functional endometrial tissue is responsive to hormonal changes and undergoes cyclic bleeding resulting in clinical and imaging findings.\(^{27}\) Adenomyosis is characterized by focal or diffuse thickening of junctional zone (>12 mm), presence of tiny T1, T2 hyperintense microcysts embedded in this thickening corresponding to islands of ectopic endometrial tissue and dilated glands, loss of junctional zone-outer myometrial interface, and adenomyoma which is focal consolidation of adenomyotic glands located within the myometrium.\(^{28,29}\) Adenomyoma may be seen as polypoidal mass protruding into uterine/cervical canal or as intramyometrial or subserosal mass. Other findings include globular enlargement of uterus, hyperintense linear striations radiating from endometrium to myometrium resulting in pseudowidening of endometrium, and adenomyotic cyst which is characterized by a hemorrhagic cyst surrounded by a solid wall. Adenomyosis has propensity to involve uterine fundus and is extremely rare in cervix.\(^{30}\) It is often seen as extension of adenomyosis from uterine fundus, body into cervix (►Figs. 17 and 18).

Isolated cervical endometriosis is considered rare. The incidence of cervix endometriosis has been reported to be between 1.6 and 2.4%, and it is often a retrospective diagnosis on histopathology. It occurs usually in women of reproductive age group; however, Park et al reported a case of cervical endometriosis in a postmenopausal woman without hormonal therapy.\(^{31}\) It may be asymptomatic or presents with postcoital bleeding/intermenstrual bleeding, rarely with severe vaginal hemorrhage.\(^{32}\) Cervical endometriosis may occur in patients without any previous history of any traumatic cervical procedures like vaginal delivery, curettage, cone biopsy, loop excision procedure, electrocautery, or cryotherapy.\(^{33}\) In a case series by Wang et al, 84.8% patients with cervical endometrosis had either vaginal deliveries or curettage procedure.\(^{34}\) Cervical endometriosis may cause abnormal smear mimicking precancerous or cancerous lesions.\(^{35}\) Sometimes cervical endometriosis may present with cystic or polypoidal cervical mass.\(^{36}\) Endometriotic deposits may involve the serosal surface of uterus extending to cervix. There may be associated involvement of broad and round ligaments which appear thickened and nodular on imaging. Deep endometriotic lesions may involve the retrocervical region (above attachment of vaginal cuff to posterior cervix), rectovaginal septum, uterosacral ligaments, posterior vaginal fornix, and anterior rectal wall. On MRI, these regions show an ill-defined soft-tissue thickening having T2-hypointense signal. Endometriotic foci show T1-hyperintense foci, and solid glandular tissue shows variable enhancement on postcontrast images.\(^{36}\)
Cervical tumors originate at squamocolumnar junction which in younger women exists at portio vaginalis, hence cancers tend to occur at portio vaginalis with exophytic growth. In older women, squamocolumnar junction exists at cervical canal and hence the tumors arise in the cervical canal, are invasive, and spread externally to involve the cervical stroma and then parametria. There is often obstruction of cervical canal with resultant hydro-/hematometra. Histologically, squamous cell carcinoma accounts for 75 to 90% of all cervical cancers. Adenocarcinoma, adenoid cystic, adenoid basal, and small-cell carcinoma form a remainder of cervical cancers. These tumors typically appear hypointense on T1W, hyperintense on T2W images, and show restricted diffusion with high signal on DWI and low signal on ADC maps. Carcinoma of cervix usually spreads to parametrium and to the uterine wall through the lymphatic invasion or deep myometrium invasion.

**Staging of cervical carcinomas**: the revised FIGO classification (2018) is used presently which considers use of any imaging modality and/or pathological findings in deciding the stage of tumor. Earlier FIGO classification (2009) was based only on clinical assessment which is observer dependent with limitations in correctly detecting parametral involvement/pelvic side wall invasion and size of tumor. Lymph node status also was not included in the earlier classification which is very important in staging of disease.40 Due to these fallacies, revised FIGO classification (2018) was made.41 The modifications are as follows:

- **Stage IA**: lateral extension measurements are removed.
- **Stage IB**: now has three subdivisions instead two.
- **Stage III**: it also has three subdivisions with IIIC indicating involvement of pelvic or retroperitoneal adenopathy which are diagnosed on radiology (r) or pathology (p). The notations “r” and “p” are to be recorded.
- **Stage I**: tumor is confined to cervix.
  - **Stage IA**: microscopic disease which cannot be evaluated on MRI.
    - IA1: Microscopic disease invasion <3 mm in depth.
    - IA2: Microscopic disease invasion >3 mm and ≤5 mm in depth
  - **Stage IB**: invasive carcinoma with measured deepest invasion ≥5 mm (►Figs. 20 and 21)
    - IB1: invasive carcinoma ≥5 mm depth of stromal invasion and <2 cm in greatest dimension.
    - IB2: invasive carcinoma ≥2 cm and ≤4 cm in greatest dimension.
    - IB3: invasive carcinoma ≥4 cm in greatest dimension.

Clinically visible lesion confined to cervix. On MRI, these tumors appear hyperintense relative to cervical stroma. The T2-hypointense rim of cervical stroma is maintained. When the thickness of this low signal intensity rim is greater than 3 mm, the hypointense rim sign is very specific (96–99%) for excluding parametral invasion thus the lesion is called definitive IB.42
Size is important in patient management as few centers offer chemoradiation for bulky tumors instead of surgery.

- Stage II: invasion beyond uterus but not to pelvic sidewalls or lower third vagina.

- Stage IIA: upper two-thirds of vagina is involved; however, no parametrial invasion. On MRI, low signal intensity vaginal wall or fornix is disrupted and is contiguous with cervical tumor (Figs. 22 and 23).
  - IIA1: tumor <4 cm.
  - IIA2: tumor ≥4 cm.

- Stage IIB: tumor invades the parametrium without reaching up to pelvic wall (Fig. 24).

On MRI, the outer T2-hypointense cervical stroma is disrupted with tumor extending into parametrium. Other findings include spiculated tumor-parametrial interface, soft tissue invading the cardinal, and uterosacral ligaments, encasement of parietal vessels and pulling of cervix to the side of invaded parametrium. In tumors with such full-thickness stromal invasion, but no parametrial mass, parametrial invasion is present in 40 to 73%.[2] However, occasionally in large tumors, it may be difficult to differentiate peritumoural edema from actual parametrial invasion. Presence of parametrial invasion is contraindication to surgery.

- Stage III: tumor extends to pelvic sidewall and/or involves the lower third of vagina and/or causes hydronephrosis/non-functioning kidney and/or involves pelvic/para-aortic nodes.

- Stage IIIA: lower third of vagina is involved without pelvic wall invasion. On MRI, there is loss of lower signal intensity wall of lower vagina (Figs. 25A and B).

- Stage IIIB: tumor involves pelvic sidewalls or causes hydronephrosis/nonfunctioning kidney.

On MRI, the tumor infiltrates the cardinal ligaments and involves the pelvic muscles. Presence of tumor within 3 mm of internal obturator, levator ani, and pyriformis muscles or the iliac vessels. Infiltration into distal ureters with proximal hydroureteronephrosis may be seen.

- Stage IIIC: involvement of pelvic and/or paraaortic lymph nodes, irrespective of tumor size and extent (with "r" and "p" notations; Fig. 26).
  - IIIC1: pelvic lymph node metastasis only.
  - IIIC2: para-aortic lymph node metastasis.

- Stage IV: tumor extends outside true pelvis or invades bladder or rectal mucosa.

- Stage IVB: distant metastases. Tumor involving organs outside true pelvis including paraaortic or inguinal nodes. Hematogenous spread of disease can occur to lung, liver, and bones.
Lymph Node Evaluation

The cervical carcinoma first spreads to pelvic nodes, that is, parametrial then obturator, internal and external nodes, and common iliac nodes. Screening of abdomen is always done in patients with carcinoma cervix to evaluate for para-aortic/retroperitoneal nodes, liver metastases.

Risk of nodal metastases increases with tumor size, depth of stromal invasion, lymphovascular invasion, and parametrial disease. MRI helps in assessment of size and morphology of the nodes. A length of 10 mm is considered to be the upper limit for short axis of normal nodes and 8 mm for rounded nodes. The diseased nodes have short-axis diameter of >10 mm, round shape, central necrosis, and perinodal infiltration. Necrosis is seen as small areas of T2-hyperintense signal in node. MRI and computed tomography (CT) have comparable overall accuracies in detecting nodal metastases; however, they cannot detect micrometastases in normal sized nodes and reactive enlargement without malignant involvement. Positron emission tomography (PET)-CT is more sensitive than CT alone or MRI and can also detect metastases in normal sized lymph nodes.

The treatment of carcinoma cervix includes either surgery and/or radical radiotherapy with adjuvant chemotherapy depending on its stage.

Cervical stage IB1 or below can be treated with surgery or definitive radiotherapy. Advantages with primary surgery are preservation of ovaries avoiding early menopause,
comorbidities, and to patients with extensive vaginal invasion that are predicted to experience severe urinary incontinence following surgery. Also, definitive radiotherapy is preferred in patients who are likely to require postoperative radiotherapy to avoid dual treatment-related morbidities.

Patients with stage IA1 may be treated with cervical conization or total hysterectomy/simple hysterectomy/extrafascial hysterectomy. As the rate of pelvic nodal metastases is less than 1%, lymphadenectomy is not recommended. However, if there is lymphovascular space invasion (LVSI), pelvic lymphadenectomy is considered with modified radical hysterectomy.55

Patients with stage IA2 are treated with modified radical hysterectomy and bilateral pelvic lymphadenectomy. Modified radical hysterectomy includes removal of uterus, cervix, medial parametrium, upper 1 to 2 cm of vagina, medial halves of uterosacral, and cardinal ligaments.55

Patients with stage IB1 are treated with radical hysterectomy or modified radical hysterectomy and pelvic lymphadenectomy. Radical hysterectomy includes removal of uterus, cervix, upper one-third of vagina, most of parametrial tissue up to pelvic wall, and removal as far as possible from the uterosacral ligaments.53

In a subset of patients with low-risk early-stage (IA2 and IB1) cervical carcinoma, rate of parametrial invasion is less than 1%. Less radical surgical options of simple hysterectomy, simple trachelectomy, cervical conization with or without sentinel node biopsy, and pelvic nodal dissection may be considered in these patients. Low-risk early-stage cervical carcinoma criteria includes squamous carcinoma, adenocarcinoma, or adenosquamous carcinoma, tumor size <2 cm, stromal invasion <10 mm, and no lymph-vascular space invasion. However, conservative approach should be considered in setting of clinical trials and only after accurate information on histology, grade, depth of invasion, margin status, and lymphovascular space invasion is assessed.54

In stages IB2 and IIA1, either surgery or radiotherapy can be used for definitive treatment. For stages IB3 and IIA2 lesions, as there are higher chances of recurrence, surgery followed by adjuvant radiotherapy is recommended. Neoadjuvant chemotherapy followed by debulking surgery is another option in bulky early stage tumors (IB3 and IIA2).
Early-stage cervical carcinoma (tumors < stage IIB) can also be treated with external beam radiotherapy plus brachytherapy. For advanced cervical carcinoma (IIB and more), chemoradiation is the optimal treatment. Surgery does not offer any additional survival benefit in these patients, increases the morbidity, and also delays radiotherapy.\(^5^5\)

**Recurrence**

Tumor recurrence can be local involving the cervix or vaginal vault, regional involving the parametrium, pelvic side walls, pelvic nodes, rectum, urinary bladder or distant involving lung, liver, para-aortic, and inguinal lymph nodes.\(^5^6\) MRI due to higher soft-tissue resolution is helpful in determining recurrence versus posttreatment change. The recurrent mass appears lobulated, T2 hyperintense, and shows restricted diffusion with heterogeneous enhancement, whereas fibrotic scar shows low signal on T2W and diffusion images\(^5^6\) (►Fig. 29). Posttreatment, the cervix becomes fibrotic and contracted along with restoration of normal low signal on T2W images, suggesting complete response to radiotherapy. The vesicovaginal and entero vaginal fistulas are most common fistulas associated with cervical cancer. These can occur due to recurrence or due to treatment related (postsurgery or postradiotherapy). MRI is useful in such cases, as it depicts exact location of fistula, single or multiple tracts, and associated with recurrence or fibrosis or pelvic adhesions. CT scan or PET-CT can help in detecting distant metastases.\(^5^7\)

**Fertility Preserving Surgery**

In patients with smaller tumors at early stage (stage I) and who desire to preservation of fertility, cervical conization or radical local excision of cervical tumor (trachelectomy) with pelvic lymphadenectomy can be performed. Radical trachelectomy includes radical local excision of cervix with adjacent parametria, laparoscopic pelvic nodal dissection, and reanastomosing the uterine isthmus to proximal vagina.\(^5^8\) Radical trachelectomy can be performed vaginally or transabdominally\(^5^7\) (►Table 3).

MRI, with sensitivity of 100% and specificity of 96%, helps in assessing the distance of tumor from the internal os.\(^5^9\) These patients also need follow-up MRI scans to detect early recurrence of tumor. The most common MRI finding post trachelectomy is uterovaginal anastomosis, posterior puckering of the vaginal wall representing “neo-fornix.” Other findings include diffuse vaginal wall thickening/edema present up to 1 year postsurgery that slowly resolve without treatment, resolving vaginal wall hematomas and submucosal fluid collections.\(^5^7\) Posttrachelectomy complications include uterovaginal stenosis, infections/abscess, and lymphoceles.

**Adenoma Malignum**

It is a subtype of mucinous adenocarcinoma of cervix with very low prevalence. Watery discharge is the initial symptom, later menorrhagia may be present. It is difficult to diagnose as imaging appearance mimics benign multiple nabothian cysts or endocervical hyperplasia. On MRI, it appears as multicystic lesion seen as T2W hyperintense signal, extending from endocervical mucosa to deep cervical stroma with enhancing solid component.\(^5^8\) It is often associated with Peutz-Jeghers syndrome.

**Extension from Endometrial Carcinoma**

Endometrial carcinoma is the most common gynecologic cancer in industrialized countries. Stage-I endometrial carcinomas are tumors confined to endometrial cavity or invading inner half of myometrium. Involvement of the endocervical glands and mucosa without cervical stromal invasion is also considered stage I. On MRI, the endometrial tumor appears isointense relative to hypointense normal endometrium on T1W images, heterogeneous intermediate relative to hyperintense to normal endometrium and hyperintense to myometrium on T2W images. Endometrial tumors enhance early on DCE and show restricted diffusion as compared with normal endometrium. These tumors are hypoenhancing as compared with myometrial enhancement. Stage-II endometrial carcinoma represents cervical stromal invasion. Stage-II cancer is associated with a higher risk of lymphovascular invasion; hence, it is important to note cervical stromal invasion on MRI\(^6^0\) (►Fig. 30). In patients with bulky uterine masses, often biopsy and histology may not establish the origin of tumor cervical or endometrial; however, immunohistochemistry may provide the diagnosis. Bourgioti et al designed a 7-point MRI scoring system to determine the origin of uterine carcinoma (cervical vs. endometrial) in histologically indeterminate cases. These are tumor location (uterine body vs. cervix), perfusion pattern, that is, tumor hypervascularity on early arterial DCE MRI, rim enhancement, depth of myometrial invasion, cervical stromal invasion, intracavitary mass, and retained endometrial secretions.\(^6^1\)

**Conclusion**

MRI is the imaging modality of choice for evaluating cervical pathologies, differentiating benign versus malignant lesions, accurately staging the malignancy, to evaluate response to treatment, detection of early tumor recurrence, and potential complications. Interpretation of posttreatment scan in patients with cervical cancer can be challenging because of expected changes in the pelvic organs. The radiologist must be familiar with these changes to avoid potential pitfalls.
Financial Support and Sponsorship

None.

Conflicts of Interest

There are no conflicts of interest.

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MRI of Uterine Cervix

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