

Primary vaginal squamous cell carcinoma arising in a squamous inclusion cyst: Case report

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SUMMARY

We report the case of a 39-year-old female with primary vaginal squamous cell carcinoma arising from a squamous inclusion cyst of the posterior wall. The tumor was located in the vaginal wall and extended into the rectovaginal septum. The overlying mucosa was intact. Histologically, there was invagination of the surface squamous vaginal epithelium forming a cystic lesion. In some areas of this invagination, the squamous epithelium showed dysplastic changes (VAIN3) transitioning into invasive squamous cell carcinoma. To the best of our knowledge, we have documented the first case of primary squamous cell carcinoma arising in a vaginal cyst in a patient without having undergone a previous hysterectomy.

Keywords: squamous cell carcinoma – vagina – squamous inclusion cyst – embryonal remnants – rectovaginal septum

Primární dlaždicobuněčný karcinom vagíny vzniklý na podkladě skvamózní inkluzní cysty zadní stěny vagíny – popis případu

SOUHRN

Prezentujeme případ 39 leté ženy s primárním vaginálním dlaždicobuněčným karcinomem vzniklým na podkladě skvamózní inkluzní cysty zadní stěny vagíny. Nádor byl lokalizován ve stěně vagíny a šířil se do rektovaginálního septa. Sliznice nad nádorem byla intaktní. Mikroskopicky byla zastižena invaginace povrchového dlaždicového epitelu, která tvořila cystickou lézi. V některých místech invaginace byly v epitelu zastiženy dysplastické změny (VAIN3) s přechodem v invazivní dlaždicobuněčný karcinom. Prezentujeme první případ primárního dlaždicobuněčného karcinomu vyrůstajícího ve vaginální cystě u pacientky, která neprodělala hysterektomii.

Klíčová slova: dlaždicobuněčný karcinom vagíny – skvamózní inkluzní cista – embryonální zbytky – rektovaginální septum

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Vaginal squamous cell carcinomas (SCCs) are rare tumors, accounting for approximately 1–2 % of all malignant neoplasms of the female genital tract (1,2). The risk factor of vaginal carcinoma is similar to that of cervical cancer, which includes a strong association with persistent human papillomavirus infection. Painless vaginal bleeding is the most common symptom of vaginal cancer. Other symptoms frequently reported include increased vaginal discharge, dyspareunia and postcoital bleeding. Most tumors occur in the upper third of the vagina and are located on the posterior wall. We report a patient with primary vaginal SCC arising from a squamous inclusion cyst of the posterior wall.

CASE REPORT

A 39-year-old woman, 2-para with a short history of dyspareunia and postcoital bleeding was referred with a palpable vaginal

tumor to the Oncogynecological center. The patient had not yet been treated for any malignancy, and had had no history of pelvic irradiation. Her last Papanicolaou smear screening, which had been done six months prior to her referral, was normal. The initial vaginal colposcopy showed the intact vaginal mucosa without any abnormality. Colonoscopic and rectoscopic examination was normal too. However, the transrectal and endovaginal ultrasound, as well as the MRI and PET, revealed a tumor, suspected to be malignant, localised in the rectovaginal septum. The spread of disease was not apparent, and both tumor markers (CA-125 and SCCA) were negative. An ultrasound-guided tru-cut biopsy from the rectovaginal septum showed a moderately differentiated nonkeratinizing invasive SCC. Subsequently, the patient underwent a radical hysterectomy type C2, partial colpectomy and the resection of the rectosigmoid bowel with terminal colostomy. No adjuvant treatment was indicated. The patient has been under regular follow-up controls every three months for 9 months without any sign of disease recurrence.

MATERIALS AND METHODS

Sections from formalin-fixed, paraffin-embedded tissue blocks were stained with hematoxylin-eosin. Selected sections were analysed immunohistochemically using the avidin-biotin complex method with the antibody directed against p16 (ready to use, Dako).

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RESULTS

Grossly, the vaginal, cervical and rectal mucosa was intact. Under the mucosa of the posterior wall of the vagina there was a firm, poorly circumscribed tumor, measuring 30 x 30 x 25 mm, extending into the rectovaginal septum (Fig. 1).

Microscopically, the uterine cervix was covered by stratified squamous epithelium. There was squamous cell metaplasia of the transformation zone. Dysplastic changes in the cervical epithelium were not found. The vagina was covered with nonkeratinizing squamous epithelium without atypia. In one area of the posterior wall, we found the invagination of the surface epithelium forming a squamous inclusion cyst (Fig. 2). In some areas of the squamous cyst,



Fig. 1. Tumor of the posterior wall of vagina.

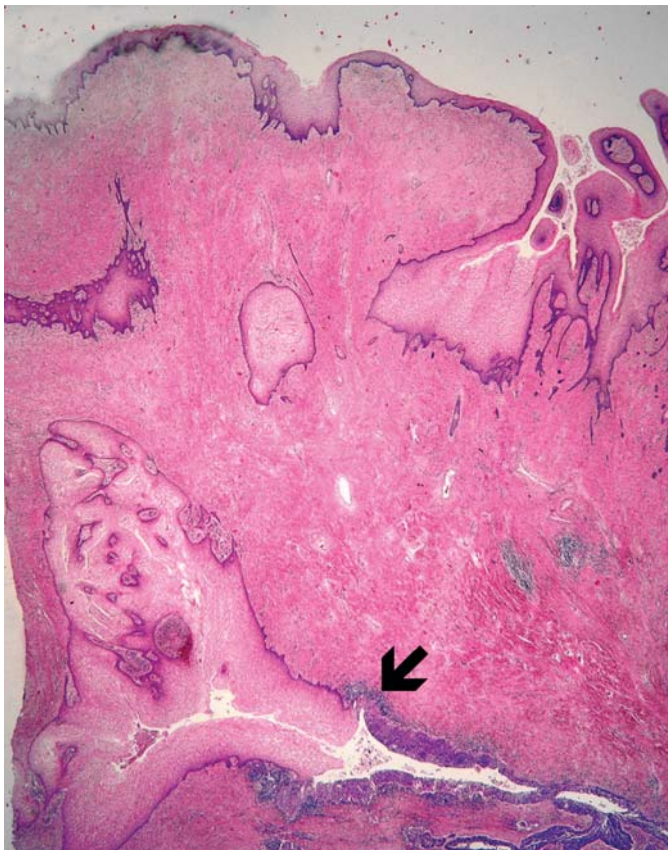


Fig. 2. Invagination of the surface epithelium forming a squamous inclusion cyst (H&E, original magnification 20x). Arrow indicates transition between normal and dysplastic squamous epithelium.

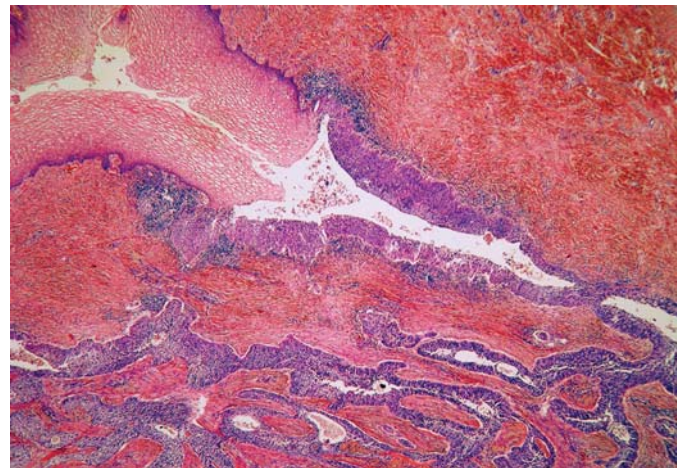


Fig. 3. Normal squamous epithelium (upper left) and dysplastic epithelium lining invagination with apparent transition into invasive squamous cell carcinoma (H&E, original magnification 100x).

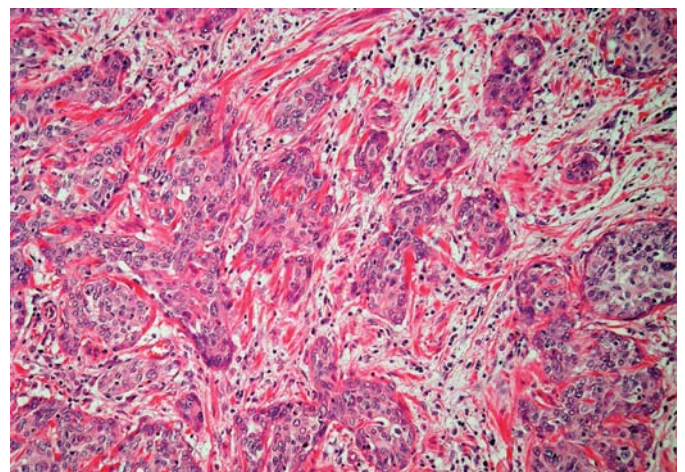


Fig. 4. Invasive squamous cell carcinoma (H&E, original magnification 200x).

the epithelium showed vaginal intraepithelial neoplasia (VAIN3) transitioning into invasive SCC with focal pseudoglandular arrangement (Fig. 3, 4). The carcinoma spread through the vaginal wall to the rectovaginal septum and the external layers of muscularis mucosae of the rectum. The rectal mucosa was intact. Immunohistochemically, the tumor cells showed a diffuse strong expression of p63 and p16.

DISCUSSION

Primary SCC of the vagina is a rare malignancy approximately accounting for 1–2 % of all malignant tumours of the female genital tract (1,2). Most vaginal SCCs represent a direct extension of either primary cervical or vulvar carcinoma. Only tumors located in the vagina not having clinical or histological evidence of involvement of the cervix or vulva can be classified as primary vaginal (1). According to the International Federation of Gynaecology and Obstetrics (FIGO), a vaginal tumor involving the uterine cervix or vulva should be classified as a primary cervical or vulvar cancer, respectively (2). Most primary tumors occur in the upper third of the vagina and are located on the posterior wall.

Cysts of the vagina are relatively uncommon. These cysts include squamous inclusion cysts, mesonephric cysts, and müllerian cysts

(1,3,4). Squamous inclusion cysts are probably the most common vaginal cysts, resulting primarily from the entrapment of fragments of mucosa during the repair of a vaginal laceration, episiotomy, or appear in vaginal cuffs after a hysterectomy. They are often asymptomatic and vary from a few millimeters to several centimeters in diameter. In rare cases, VAIN or invasive SCC can arise in vaginal squamous inclusion cysts. These instances are rare, and only three cases of VAIN and 2 cases of SCCs arising within vaginal cuff inclusion cysts or sinuses have been described in patients who had previously undergone a hysterectomy (5). In one of these patients, the hysterectomy was performed because of cervical cancer, in two for cervical intraepithelial neoplasia, and in the other two the cause for hysterectomy was not reported. In our case, however, the tumor arose from the invagination of squamous epithelium located in the posterior vaginal wall, which most likely represented a squamous inclusion cyst in a patient without a previous hysterectomy. The patient had neither an anamnestic nor current clinical or histological evidence of the involvement of the cervix or vulva.

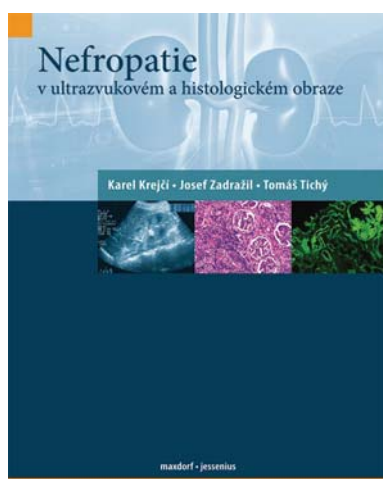
Differential diagnosis in our case includes the origin of SCC from embryonal remnants in the rectovaginal septum or in the longitudinal vaginal septum, or from endometriosis. However, the

tumor origin from the squamous inclusion cyst of the posterior wall is far more likely, because we had found the particular site of the invagination of the surface vaginal epithelium. Tumors arising from endometriosis of the rectovaginal septum are rare neoplasms, which can mimic either primary vaginal or rectal tumors. However, these tumors mostly represent carcinomas of müllerian origin such as endometrioid and serous adenocarcinoma. Moreover, the foci of endometriosis are usually found close to the tumor (6).

In conclusion, we described a case of primary SCC probably arising from a squamous inclusion cyst of the vaginal wall. To the best of our knowledge our case represents the first case of SCC arising in a vaginal cyst in a patient without a prior hysterectomy. Only two other cases of SCCs arising in vaginal squamous cysts have been described to date, however, these tumors were found in patients who had previously undergone a hysterectomy (5). In addition, one report described a case of vaginal SCC which was cystic and grew endophytically into the rectovaginal septum (7). The possibility of carcinomas originating in a vaginal wall cyst should be borne in mind in the differential diagnosis of tumors located in the vaginal wall and the rectovaginal septum.

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NEFROPATIE V ULTRAZVUKOVÉM A HISTOLOGICKÉM OBRAZE

RECENZE

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Monografie „Nefropatie v ultrazvukovém a histologickém obraze“ vyplňuje mezeru v českých publikacích zabývajících se problematikou ultrasonografie v nefrologii a nefropatologii. Shrnuje mnohaleté zkušenosti autorů získané při hodnocení ultrazvukových a histologických obrazů v diferenciální diagnostice nefropatií autologických i transplantovaných ledvin a monitorování potransplantačních komplikací. Kniha využívá také závěrů vědeckovýzkumné činnosti a přináší řadu nových poznatků, které byly autory prezentovány jak v našich, tak i v zahraničních periodikách. Podnětem k jejímu vydání byly rovněž zkušenosti získané při pořádání sonografických kurzů pro nefrology a internisty v rámci Institutu postgraduálního vzdělávání ve zdravotnictví. Atlas je určen k dalšímu vzdělávání a studiu především pro nefrology, urology, pracovníky transplantáčnických center, internisty provádějící ultrazvuková vyšetření ledvin, radiology a klinické patology zabývající se nefropatologií. Může být cennou pomůckou i pro studenty medicíny s hlubším zájmem o problematiku nefrologie a ultrasonografie.