Paraganglioma of the vagina: a case report and review of the literature

Abstract: Vaginal paraganglioma is a rare and unusual tumor occurring in the vaginal wall. It is a solitary primary paraganglioma, especially in atypical sites. Herein, we report an unusual case of a 17-year-old woman who had not experienced vomiting, or hypertension. She was found to have an immobile solid mass in the right side of her vaginal wall. Positron emission tomography/computed tomography scans revealed a well-defined solid ovoid mass adjacent to the bladder and pelvic floor. Tumor markers were within the normal range. A transient blood pressure increase occurred during the biopsy. After oral administration of antihypertensive drugs, surgery was performed to completely remove the mass. Histopathological examination indicated that it was a paraganglioma of the vagina. Repeat computed tomography examination did not reveal any local recurrence or distant metastasis during the 12-month follow-up period.

Keywords: vaginal paraganglioma, neuroendocrine tumor, clinical features, treatment

Introduction

Derived from neural crest cells, paraganglioma is formed by chromaffin or nonchromaffin neural epithelioid cells, which are related to sympathetic or parasympathetic nerves.1 Until now, there has been no uniform standard for its phase division and treatment.2 Paragangliomas are mostly distributed in paraganglion-gathering areas, such as the adrenal glands, neck, mediastinum, or retroperitoneum,3–5 but rarely occur in the vaginal wall. This report will analyze one case of a young woman who suffered from vaginal paraganglioma, together with a review of other relevant literature.

Case report

A 17-year-old woman with no history of marriage or pregnancy was hospitalized because of irregular vaginal bleeding over 5 months. She had no history of genetic disease, coagulation dysfunction, or liver cirrhosis, but had asthma. Physical examination revealed a blood pressure of 140/100 mmHg, a pulse rate of 95/minute, and no superficial lymph nodes could be palpated. Since the patient had had no sex, abdomen ultrasound was used instead of vaginal ultrasound. This showed that her uterus appeared normal in its dimensions, shape, and position. Both ovaries seemed normal for site, dimension, and echo structure. There was no free fluid in the Douglas pouch, while her pelvic computed tomography (CT) showed a 40 × 32 mm regular-margin hyperdense abnormal signal in the right vagina wall adjacent to the bladder. The results of positron emission tomography (PET)/CT were as follows: mass of soft-tissue density in the right vaginal wall and unusual hypermetabolism of fluorodeoxyglucose (FDG) matched with the tumor appearance; nodular soft-tissue density shadow in the pelvic wall.
cavity and hypermetabolism of FDG may have indicated the same lesion in the right vaginal wall. During biopsy of the vaginal tumor under intravenous anesthesia, her blood pressure suddenly rose to 160/100 mmHg, and then came back to 130/90 mmHg after examination. The pathological diagnosis was vaginal paraganglioma.

After oral administration of sustained-release metoprolol and nifedipine tablets for 1 week, the woman had vaginal tumorectomy and reconstruction performed, and she had smooth blood pressure throughout the operation. The tumor mass was gray-white and taupe, with a size of 35×30×25 mm (Figure 1). Microscopically, polygon-like oncocytes were spread in nest or cluster forms. Abundant acidophilic granulates within the cytoplasm and round or oval karyons could be clearly identified. The tumor cells were clustered in small nests or irregular trabecular structures demarcated by delicate fibrous stroma and capillaries in the Zellballen pattern. In the tumor stroma, blood sinus, blood capillary, and little fibrous tissue could be found (Figure 2). Synaptophysin, chromogranin A, protein S-100, neuron-specific enolase, cell proliferation-associated antigen Ki67, and vimentin were all positive, while smooth-muscle actin, human melanoma-associated antigen (HMB-45), and cytokeratin were negative in immunohistochemistry assay (Figure 3).

**Discussion**

Paraganglioma is a rare soft-tissue neuroendocrine tumor, and can be classified as pheochromocytoma and extra-adrenal paraganglioma depending on its pathogenic site. The latter is closely related to tumors in the extra-adrenal sympathetic and parasympathetic paraganglioma, and named after the anatomic site as well as its functional activity. Paragangliomas have rarely been reported in the female genital tract, including in the ovary, broad ligament, uterus, and vulva. To the best of our knowledge, only five cases of vaginal paraganglioma have been reported since 1998, with a median age of 31. 3 years if including our case together.

Furthermore, paragangliomas can be divided into functional and nonfunctional types according to their ability for synthesis, storage, and secretion of catecholamine, as well as production of various peptides, neurohormones, and chromaffin protein grains. Paraganglioma is typical in its clinical manifestations, such as paroxysmal or consecutive elevation of blood pressure, headache, dizziness, palpitation, sweating, fainting, and blurred vision. It is also known as catecholamine syndrome. Located in the vagina, paraganglioma has some special symptoms, such as irregular vaginal bleeding, vaginal pain, or finding a vaginal wall mass during routine gynecological examination. Sometimes, serious cardiovascular complications like hypertension, tachycardia, arrhythmia, or myocardial infarction may arise from the stimulation of surgery or coitus. In this case, the young woman had irregular vaginal bleeding, and suffered from dizziness, headache, and palpitation after physical exercise. Her blood
pressure and heart rate fluctuated abnormally during the biopsy examination, which coincided with functional nonchromaffin paraganglioma.

It is difficult to make a preoperative diagnosis of vaginal paraganglioma due to the lack of specific symptoms, and it may be misdiagnosed as vaginal rhabdomyosarcoma, leiomyoma, or hemangioma, among others. Therefore, it should be definitively diagnosed based only on histological findings.22

Ultrasound, CT, or magnetic resonance imaging is useful for delineating the location, while 18F-FDG-PET helps to estimate malignant, multiple, and metastatic lesions.23,24 In this case, the vaginal mass was not found by ultrasound, probably because it was not examined by transvaginal ultrasound, although it unusual hypermetabolism of FDG under PET/CT appeared, which matched with the tumor’s features. It has been reported that 123I-metaiodobenzylguanidine has strong affinity to chromaffin cell tissues when used as tracer imaging to effectively diagnose or locate paraganglioma. However, this detection technology had not been widely used.25 Moreover, an assay of catecholamine and noradrenaline in blood and urine also helped to diagnosis paraganglioma. Patients with functional nonchromaffin paraganglioma may have an increase of catecholamine and noradrenaline in blood and urine, along with blood pressure elevation.26 Such information was not available in this case.

Whenever it comes to either functional or nonfunctional vaginal paraganglioma, all the literature suggests that surgical operation currently seems to be the most effective treatment. Because it cannot be completely determined as functional vaginal paraganglioma before operation, the perioperative period is very important.27 Usually, an α-receptor antagonist is used for catecholamine inhibition. It has been reported that interventional embolization of uterine artery was carried out in one case.18 During the operation of the uterine artery, fluctuation of blood pressure, arrhythmia, pneumoniedema, myocardial ischemia, or shock may be life-threatening. Those complications can be decreased to 3% by sufficient preoperative preparation.

It is widely recognized that paraganglioma is a slowly growing tumor, with no or low latent malignancy. However, it cannot be determined just by histological morphology. Instead, it should be judged malignant on the bases of neoplasm recurrence, metastasis of lymph nodes or nonchromaffin tissues, and invasion or infiltration to the adjacent apparatus.29–31 Due to the possibility of neoplasm recurrence and metastasis, as reported, patients need periodic checks and long-term follow-up.27–31 In this case, the woman had complete resection of the tumor as well as vaginal reconstruction performed. Repeat CT examination did not reveal any local recurrence or distant metastasis during the 12-month follow-up period. We suggested the patient have a yearly pelvic CT scan and gynecological examination.

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Disclosure
The authors report no conflicts of interest in this work.

References