Surgical Management of an Extragonadal Trabecular Carcinoid Tumor: A Case Report and Review of the Literature

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Abstract
Extragonadal malignant transformation of a teratoma is rare, with only six reported cases in the literature. We report a case of a 33-year-old gravida 4 para 2 woman who had incidentally discovered bilateral pelvic masses, which were followed radiographically for nine years. She then underwent surgical resection, which revealed metastatic trabecular carcinoid involving lymph node tissue (left retroperitoneal pelvic mass), and trabecular carcinoid arising in a background of mature teratoma (right presacral mass). This case illustrates the indolent nature of this tumor and the importance of complete surgical resection.

Keywords: Extragonadal germ cell tumors; Trabecular carcinoid tumors; Mature teratoma

Introduction
Malignant extragonadal germ cell tumors can arise anywhere along the gonadal ridge. Locations may include the retroperitoneal, mediastinum, presacral region, and pineal gland [1]. An Extragonadal Germ Cell Tumor (EGCT) is by definition a germ cell neoplasm displaying one of the histologies associated with gonadal origin, but located outside of the gonads [2]. EGCT arise from primordial germ cells that migrate from the yolk sac/endoenderm to the genital ridge [3]. The standard treatment for EGCT includes surgery, while chemotherapy or radiotherapy may be added for certain malignant subtypes [4].

Carcinoid tumors are uncommon neoplasms in the diffuse peripheral endocrine system. Ovarian carcinoids are uncommon and primary ovarian carcinoids make up approximately 0.5-1.7% of all carcinoid tumors [5]. Extragonadal carcinoid tumors are extremely rare with only six reported cases in the literature [6-11]. We report on a patient with a slowly growing pelvic mass observed radiographically over nine years that at surgical excision was found to be an extragonadal trabecular carcinoid.

Case Report
33-year-old Caucasian female patient, gravida 4 para 2, without other significant past medical history presented in 2014 after the birth of her second child for management of stable bilateral pelvic masses that had been monitored since 2005. Lesions were initially noted as an incidental finding on a CT of abdomen and pelvis performed in the setting of appendicitis in 2005. They were not noted or commented upon in the operative note from a laparoscopic appendectomy performed at that time. They were then re-identified on routine obstetric ultrasound, and were conservatively managed with watchful waiting in the setting of her pregnancy. She had an uncomplicated pregnancy and underwent a primary cesarean section for malpresentation. The retroperitoneal masses were not commented on in the operative report.

Postpartum, she was referred for definitive surgical management and diagnosis. The patient was entirely asymptomatic. She had no irregular bleeding, and no lower extremity or back pain. She experienced constipation and irregular bowel movements for several years. Physical exam was unremarkable, with normal lower extremity motor and sensory nervous systems, and a pelvic exam without obvious palpable adnexal masses.

She underwent a follow-up pelvic MRI in December of 2013 that showed a slight increase in the size of the left sided mass. She then underwent surgical excision in February of 2014. Figure 1 shows the clinical timeline.

Radiology
Table 1 outlines the radiologic imaging studies. Figure 2 shows ultrasound and CT scan images from 2012. Figure 3 shows the July 2013 axial MRI images of the retroperitoneal position of the masses. Figure 4 shows a sagittal MRI image of the presacral mass from December of 2013.

Surgical Procedure
A laparoscopic approach was used. The left retroperitoneal space was entered and a 6 cm mass was identified along the left pelvic sidewall, inferior to the ureter (Figure 5). The ureter was identified and fully mobilized and separated from the mass. The feeder vessels from the internal iliac vessels that vascularized the mass were separately isolated and cauterized. The mass was resected completely without rupture and placed in an endoscopic bag and removed.

The right sided retroperitoneal mass was adherent to the right presacral space just superiorly to the cul-de-sac and was densely adherent to the foramina of the sacrum. The mass was inferior and at a distance from the right ureter and right iliac vessels. An incision was made in the peritoneum of the medial leaf of the right broad ligament, inferior to the ureter, and the rectum was mobilized anteriorly. The mass was gently peeled off the sacral nerves and veins using a laparoscopic swab and harmonic cautery. The mass was unruptured and was placed in an endoscopic bag and removed.

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In 2005, the patient underwent a laparoscopic appendectomy for appendicitis. Pathology showed acute suppurative appendicitis and periappendicitis. There was no evidence of a carcinoid tumor within the appendix.

In 2014, the patient underwent a resection of the right presacral mass and the left retroperitoneal mass. The right presacral mass measured 3.8 by 2.6 by 2.1 cm and its cut surface showed a 2.2 × 1.5 × 1 cm smooth lined cyst containing pink gelatinous material that was surrounded by homogeneous, solid and focally lobulated tan tissue with speckles of hemorrhage (Figure 6). The left retroperitoneal mass measured 6.5 × 6.2 × 1.4 that had a tan soft appearance.

On microscopic evaluation, the left retroperitoneal pelvic mass was felt to be consistent with metastatic trabecular carcinoid involving lymph node tissue, whereas the right presacral mass was consistent with trabecular carcinoid arising in a background of mature teratoma. The cystic portion of the sacral mass was lined by benign-appearing mucinous, pseudostratified ciliated and transitional-like epithelium. Scattered islands of fetal cartilage were present (Figure 7). As mentioned, the overall appearance was that of a mature teratoma. However, most of the presacral mass and all the retroperitoneal mass were composed of a trabecular carcinoid in which irregular often anastomosing trabeculae display cells with relatively abundant eosinophilic cytoplasm and homogeneous small oval to round nuclei with “salt and pepper” chromatin (Figure 8). Mitoses were rare. The stroma was typically scant but prominent thin walled blood vessels were frequently found. Only this component of the tumor was positive for neuroendocrine markers including chromogranin, synaptophysin, NSE and CD56 (Figure 9).

Follow-up Clinical Course

In the initial postoperative period, the patient underwent a full metastatic work-up. Her CA 125 was 7 units/ml (normal range 7-35). Human chorionic gonadotropin was less than 6 mIU/ml. The twenty-four urine 5 hydroxyindole acetic acid (5HIAA) was in the normal range at 2 mg/24 hour. A PET CT scan in August of 2014 was negative for lesions and for metabolic uptake of 2-deoxy-2-[fluorine-18]fluoro-D-glucose (18F FDG).

The patient became pregnant in the summer of 2015 but had a pregnancy loss at 20 weeks. At her October 2015 follow-up, twenty-one months after the surgical resection of the extragonadal germ cell tumor and the metastases in one contralateral lymph node, the pelvic MRI was negative for masses. The patient is currently in good health and has no complications from the surgical resection.

Discussion

Extragonadal germ cell tumors have major differences in clinical behavior that suggest EGCT are biologically different from primary gonadal germ cell tumors [2]. For instance, EGCT are much more likely to be bulky at presentation and as with our patient, have metastatic disease.

In one large international review of 635 consecutive patients treated from 1975 through 1996 at 11 cancer centers, 341 patients (54%) had primary mediastinal EGCT, and 283 patients (45%) had...
Intraoperative Photograph: Left pelvic sidewall. Ureter (white arrow) has been mobilized superiorly. Retroperitoneal mass (black arrow) is inferior to the iliac vessels.

Figure 5: Intraoperative Photograph: Left pelvic sidewall.

Axial post contrast MRI images demonstrate enhancing pelvic masses in the right presacral space and the left retroperitoneum (white arrows). Two non-enhancing cystic areas are seen in the left retroperitoneal mass (bottom image).

Figure 3: Pelvic MRI from July 2013.

Sagittal MRI image demonstrates presacral mass (*), with uterus also identified (**).

Figure 4: Pelvic MRI from December 2013.

Table 1: Radiologic imaging studies from 2005 to 2014 of an extragonadal germ cell tumor.

Intraoperative Photograph: Left pelvic sidewall. Ureter (white arrow) has been mobilized superiorly. Retroperitoneal mass (black arrow) is inferior to the iliac vessels.

Figure 5: Intraoperative Photograph: Left pelvic sidewall.

Axial post contrast MRI images demonstrate enhancing pelvic masses in the right presacral space and the left retroperitoneum (white arrows). Two non-enhancing cystic areas are seen in the left retroperitoneal mass (bottom image).

Figure 3: Pelvic MRI from July 2013.

Sagittal MRI image demonstrates presacral mass (*), with uterus also identified (**).

Figure 4: Pelvic MRI from December 2013.

retroperitoneal EGCT. Five hundred twenty-four patients (83%) had a nonseminomatous germ cell tumor (GCT), and 104 patients (16%) had a seminomatous histology [12].

Malignant germ cell tumors are uncommon, 1-3% of germ cell tumors, and the majority of these malignancies are embryonal tumors such as dysgerminomas, embryonal carcinomas, or yolk sac tumors [13]. Much more rare is the malignant transformation of the somatic
components of a mature cystic teratoma, of which squamous cell cancer accounts for 70-88% [14]. Among mature cystic teratomas with malignant transformation, neuroendocrine carcinoid tumors are significantly more rare, with reports in approximately 6% of all malignant cases [15]. Only six extragonadal carcinoid tumors arising in mature cystic teratomas have been reported in the English literature to date, but none of these cases reported metastatic disease [6-11]. Table 2 summarizes the clinical findings of these individual cases. Our patient’s case is the first report of extragonadal malignant transformation with metastasis, found in lymph node tissue.

In the literature, age range varies widely, from 22 to 63 years old (mean 40.8 years), with the most common presenting symptom being abdominal pain or discomfort. Radiologic findings are similar in all cases, with complex cystic and solid tumors. The reported size of these tumors is large, ranging from 5 cm to 16 cm, with a mean of approximately 11cm in largest diameter. In the case presented here, the patient presented initially asymptomatic with a small 2 cm lesion, which then grew on follow up imaging examination to 3 cm and was noted to be 6 cm at surgery. It is possible that these lesions would not have been discovered, had they not been noted on the obstetrical ultrasounds, until they became large enough to cause symptoms.

Classically, as with our patient, these tumors are treated with surgical excision. The role of chemotherapy or radiation has not been established. Two of the six case reports included a referral to chemotherapy [6,10]. In carcinoid tumors arising from teratomas of the ovary, treatment can be completed with surgical excision alone, usually salpingo-oophorectomy [16]. There are reports of five-year survival with cystectomy alone for primary ovarian carcinoid malignancies [15]. There was only one case study that reported metastasis from the ovary of the carcinoid element, unsuccessfully treated with thioTEPA [17].

Our patient was treated by a radical retroperitoneal excision using a minimally invasive surgical approach. Given the deeply placed position of extragonadal tumors over major vascular structures and neurovascular bundles, a laparoscopic approach should be encouraged. Laparoscopy offers higher resolution optics, which allows the ability to preserve nerves and blood vessels, with a significant reduction blood loss [18].

Trabecular carcinoids may be difficult to diagnose preoperatively. Our patient had normal tumor markers and a very slow growth of the tumor over a period of 8 years. Other reports also document a lack of tumor markers in this type of tumor [5]. Carcinoid tumors are defined as of neuroendocrine origin and have been classified into 3 types: neuroendocrine carcinomas, typical and atypical carcinoids. While neuroendocrine carcinoma is a malignant epithelial neoplasm with high mitotic activity, high nucleoplasmic ratio, necrosis, vascular invasion, high rate of metastasis, and poor prognosis [19], trabecular carcinoids are much more indolent. The typical carcinoid tumor is a low-grade malignant tumor with rare mitotic activity, rare necrosis, monotonity of nuclear size and shape, slow growth, and good prognosis. The atypical carcinoid tumor is an intermediate-grade neoplasm generally with punctuated necrosis and intermediate-mitotic activity [20].

Information from large series is instructive to understand behavior and prognosis. An institutional review evaluated 26 ovarian carcinoids and determined that 16 were primary and 10 were metastatic from the gastrointestinal tract [21]. The average size of primary ovarian carcinoids was 3.4 cm (range: 0.2-13.5 cm) versus 10.2 cm for metastatic carcinoids (range: 4-32 cm). Teratomatous elements
were present in association with 10/16 primary ovarian carcinoids, whereas none were present in any metastatic carcinoid. The primary ovarian carcinoid types were insular (n=6), trabecular (n=3), strumal (n=6, of which 5 were trabecular pattern and 1 was insular pattern) or mucinous (n=1). All metastatic carcinoids, except for two of mucinous type, were insular [21].

Based on the six previously reported extragonadal carcinoid tumors, the prognosis after excision is excellent [6-11]. Disease-free survival was reported as 6 to 31 months. Our patient has no evidence of recurrence from a metastatic extragonadal carcinoid at 21 months to date.

Conclusion

1) Malignant transformation in a mature cystic teratoma to a trabecular carcinoid is uncommon, but these tumors are regarded as having a low malignant potential with good prognosis and survival.

2) Extragonadal germ cell tumors are exceedingly rare and are difficult to diagnose without surgery.

3) Complete surgical excision is the treatment of choice. Minimally invasive surgical techniques such as laparoscopic retroperitoneal dissection should be encouraged.

4) This report describes the seventh case of an extragonadal carcinoid tumor and the first case of metastatic disease in a lymph node.

References


