Pancreatic parangglioma with multiple lymph node metastases found by spectral CT: A case report and review of the literature

Li T et al. Pancreatic parangglioma with lymph node metastases

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Abstract

BACKGROUND

Primary pancreatic parangglioma is exceedingly rare. Most patients lack a typical clinical presentation, and the tumor is difficult to accurately differentiate from other pancreatic neuroendocrine tumors, making the misdiagnosis rate extremely high. Surgical excision is the primary treatment modality but is considered high risk. Because of its rich vascularity, the tumor easily bleeds during surgery, especially malignant paranggliomas invading large blood vessels. Thus, a thorough preoperative evaluation of the tumor is necessary. Here, we report a primary malignant pancreatic parangglioma, the second such case in a young patient that was successfully resected surgically.

CASE SUMMARY

A 26-year-old female patient was admitted to the hospital with unexplained abdominal pain. Dual-layer spectral-detector computed tomography (DLCT) revealed a mixed density mass in the pancreatic body tail. The patient was transferred to our hospital after previous failed surgical resection at other hospitals. The patient and family strongly desired surgery. After a thorough preoperative evaluation and adequate preparation, a large mass with a greatest dimension of 8.0 cm was successfully resected. The final pathological diagnosis was malignant parangglioma. The patient was discharged in good condition two weeks postoperatively.

CONCLUSION
The rare malignant pancreatic paraganglioma reported here was difficult to diagnose preoperatively. Early filling of the draining vein may be a crucial diagnostic imaging feature. DLCT can provide more precise information for surgical resection through dual-energy imaging.

**Key Words:** Spectral computed tomography; Pancreas; Paraganglioma; Lymph node metastasis; Case report


**Core Tip:** Primary pancreatic paraganglioma is exceedingly rare, generally benign tumor, with only 4 cases of malignancy reported. They are often misdiagnosed as pancreatic neuroendocrine tumors (pNETs). Early filling of the draining veins may be a crucial imaging feature to differentiate these tumors from other pNETs and may be observed more frequently in malignant cases. Surgical resection is the primary treatment modality. However, the rich vascularity and potential functionality of the tumor pose a significant risk for invasive surgery. Thorough preoperative evaluation and preparation are necessary. Definitive diagnosis relies primarily on histopathological examination.

**INTRODUCTION**

Paraganglioma originates from neural crest cells in the sympathetic or parasympathetic ganglia and is a rare neuroendocrine tumor with an incidence of approximately 2-8 per 1 million people per year[1]. Paragangliomas arising in the pancreas are even rarer, with an average age of onset of 52 years, and most of these tumors are nonfunctional[2]. The malignancy rate of paragangliomas is approximately 10%-50%[3]. Patients often lack a typical clinical presentation, especially those with nonfunctional paragangliomas, and
the imaging features of these tumors are similar to those of pancreatic neuroendocrine tumors (pNETs), making the preoperative diagnosis very difficult. Herein, we report a malignant pancreatic paraganglioma in a young person. We discuss the imaging features and clinical characteristics of the tumor and review the relevant literature.

CASE PRESENTATION

Chief complaints
A 26-year-old female patient complained of epigastric pain for two years.

History of present illness
This patient had frequent epigastric pain accompanied by posterior back pain starting two years prior, which could not be relieved by changing positions. A pancreatic mass was found at a local medical institution, and a frozen tissue biopsy revealed a possible rhabdomyosarcoma. The mass was assessed as unresectable, and the operation was halted. The patient was then transferred to a higher tier hospital for treatment, where the previous biopsy was retrieved and with the diagnosis of mucinous spindle cell soft tissue tumor favored. The patient was discharged after receiving chemotherapy (paclitaxel liposome + nedaplatin four times), radiotherapy (25 d), targeted therapy (anlotinib), and immunotherapy (toripalimab), which was poor treatments, and the above therapeutic measures were discontinued after more than one year.

History of past illness
The patient previously had elevated blood glucose levels up to 21 mmol/L and was not receiving regular treatment. She denied hypertension or other medical histories.

Personal and family history
She had no history of alcohol or tobacco abuse. There was no obvious abnormality in her family history.
**Physical examination**

There was no obvious abnormality in her physical examination. Her abdomen was soft, without tenderness or a palpable mass.

**Laboratory examinations**

Laboratory examinations showed that the patient's fasting glucose and antithyroid peroxidase antibody (TPO-Ab) levels were 9.03 mmol/L and 359.88 IU/mL on admission. The serum tumor markers were within the normal ranges. There were no plasma/urine levels fractioned metanephrines and catecholamines measured.

**Imaging examinations**

Abdominal dual-layer spectral-detector computed tomography (DLCT) showed a mixed density mass in the pancreatic body tail, measuring approximately 7.1 cm × 5.7 cm × 3.7 cm, with indistinct borders and short streaks of calcification within the mass (Figure 1A). The contrast-enhanced (Figure 1B) scans showed significant mass enhancement, especially in the arterial phase, and patchy hypointense areas were observed within the mass. The peritumor and intratumor vessels were abundant, and the draining veins of the mass were observed early and converged into the portal venous system (Figure 1C), with extensive tortuous and dilated veins in the portal venous basin. The standard portal vein trunk structure disappeared and showed sponge-like changes. The mass enveloped the splenic artery and splenic vein, and multiple abnormally enhanced and enlarged lymph nodes were observed in the peripancreatic fatty and hepatogastric spaces. The largest lymph node was approximately 3.4 cm, with circumferential enhancement and central necrosis visible within the lymph node. The upper edge of the mass was adherent to the gastric wall, and there was no clear invasion of the gastric wall (Figure 2). The mass (Figure 1D-E) showed low intensity on T1-weighted imaging (T1WI) and slight hyperintensity on T2 WI. Magnetic resonance cholangiopancreatography showed no dilatation of the bile
duct, hepatic ducts, or pancreatic duct (Figure 1F). The imaging diagnosis indicated a malignant neuroendocrine tumor.

**FINAL DIAGNOSIS**

Postoperative pathology uncovered a solid gray–white mass measuring approximately 8 cm × 5 cm × 4 cm. Histological examination revealed a tumor consisting of well-defined nests of polygonal cells separated by vascular fiber septa, forming a classic Zellballen pattern (Figure 4). Immunohistochemistry showed positivity for chromogranin A, synaptophysin, as well as sustentacular cells expressing S-100 protein. The Ki-67 index was 8%. Staining for cytokeratin and epithelial membrane antigen was negative. Therefore, based on a combination of the histology and immunohistochemistry results, the final diagnosis was malignant pancreatic paraganglioma, with tumor metastases in the peripancreatic lymph nodes (4/6) and no metastases in the excisional margin or spleen.

**TREATMENT**

Based on the imaging findings, the patient underwent pancreatic body-tail resection, total splenectomy, and radical lymph node dissection. A large and solid tumor originating from the pancreatic body-tail was found during surgery. The long diameter of the tumor was approximately 8 cm, the mass was adherent to the stomach's posterior wall, and the lower portion of the tumor was located in the mesentery of the transverse colon and to the left of the superior mesenteric artery. A hard lymph node of approximately 3.5 cm in diameter was found near the lesser curvature of the stomach. The patient's blood pressure levels and heart rates were not significantly altered during the procedures.

**OUTCOME AND FOLLOW-UP**

The patient was transferred to the intensive care unit after surgery and was discharged in good condition two weeks later; the patient did not receive follow-up adjuvant
therapy. The patients' blood glucose levels and TPO-Ab levels failed to return to normal.

**DISCUSSION**

Paraganglioma is a rare tumor that originates from neural crest cells. This tumor usually occurs in the head, neck, and retroperitoneum, and paragangliomas in the pancreas are rare\(^4\). Malignancy is even rarer, with only four cases reported to date (Table 1)\(^5-8\), including three patients who showed lymph node metastases and one patient with multiple liver metastases. Currently, the presence of invasion into vascular and peripheral structures or metastasis is a reliable basis for diagnosing malignant paraganglioma\(^9\). We reported a malignant nonfunctional paraganglioma in the pancreas with lymph node metastasis, the second such case in a young patient.

Patients with paragangliomas of the pancreas usually lack specific clinical manifestations. Patients with nonfunctional paragangliomas often have no obvious symptoms or present with unexplained epigastric pain, and such tumors are mostly found incidentally during imaging examinations. Due to elevated catecholamine levels, patients with functional paragangliomas may feel hypertension, headache, sweating, and palpitations\(^10\). However, these manifestations are easily overlooked by clinicians, and the presence of a functional tumor is often realized only when a rapid increase in blood pressure and heart rate occurs during surgery. Due to the mass's rich vascularity and potential functionality, blind invasive investigations and procedures may result in catastrophic complications or surgical failure. For instance, a patient\(^7\) with pancreatic paraganglioma had a sudden rise in blood pressure to 220 mmHg during surgery. The operation had to be stopped because of inadequate preoperative preparation. Another patient\(^11\) was preoperatively misdiagnosed with pancreatic cancer. After touching the tumor intraoperatively, the patient's systolic blood pressure suddenly rose 180 mmHg, and her heart rate reached 140 beats per minute. As a result, the operation was stopped, and the patient died from cardiac failure 34 h after the operation. These cases indicate that the relevant laboratory tests performed before surgery must be improved to
prevent failed surgeries and death due to functional tumors and inadequate preoperative preparation. Meta-iodobenzylguanidine (MIBG) nuclear imaging and measurement of metanephrines in blood or urine can effectively differentiate between functional and nonfunctional paragangliomas[12].

Imaging examinations are an essential method for diagnosing and evaluating pancreatic paraganglioma. The parenchymal portion of the mass shows low-intensity on T1WI and hyperintensity on T2WI[13]. CT scans usually show a solid or cystic soft tissue mass, most often located in the pancreatic head, with some calcifications visible. The mass is heterogeneously enhanced with necrosis and cystic changes in the arterial phase and persistently enhanced in the portal and venous phases, with abundant peritumoral and intratumoral vessels. Although the head of the pancreas is the most common site of pancreatic paraganglioma and the tumor is usually large, there is no significant dilatation of the bile duct and pancreatic duct. In a few cases, the pancreatic duct is mildly dilated[14-18]. Pancreatic cancer often presents as a hypoenhamcing mass in the pancreatic head with significant dilation of the pancreatic and bile ducts, which allows the differentiation of pancreatic paragangliomas from pancreatic cancer. Patients with pancreatic paragangliomas usually do not have elevated serum tumor markers. Because of the similar imaging presentations, it is difficult to distinguish paragangliomas from other types of pNETs; however, it has been reported[15,18] that paragangliomas often present with early filling of the draining veins, and in 50% of evaluable cases, the draining veins can be observed. In our case, we also found a large number of draining veins. Moreover, this sign seems to be more visible in malignant cases. Early filling of the draining veins has been observed in all evaluable malignant cases (Table 1). This sign may be a vital imaging feature for diagnosing pancreatic paraganglioma.

Although it is difficult to differentiate pancreatic paragangliomas from pNETs accurately, surgical treatment is the preferred treatment for both. Therefore, preoperative assessments of tumor size, degree of invasion, and relationship with surrounding blood vessels and adjacent organs are more important. CT is currently the
main imaging examination for pancreatic tumor evaluation. DLCT can separate X-ray photons into two energy levels during the detector readout, enabling the generation of both conventional CT images and images based on dual-energy processing. When using DLCT, typical reconstructed image sets include virtual monoenergetic images at varying energy levels, effective atomic number maps, and material-decomposition images (e.g., maps of water and iodine content). Therefore, DLCT can provide information beyond conventional CT analysis. We performed a comprehensive imaging evaluation of the lesion using DLCT. Based on the powerful advantages of dual-energy imaging and multiparametric imaging, we more accurately assessed the boundary of the lesion, the distribution of the surrounding vessels, and the presence of lymph node metastasis. Moreover, we graphed the hounsfield unit attenuation plot of the surrounding lymph nodes and mass and assessed lymph node metastasis according to the slope (Figure 3). Surgical excision is the primary treatment modality. The surgical approach for pancreatic paraganglioma depends on the tumor's location, including pancreaticoduodenectomy (PD) or pylorus-preserving PD (PPPD) for tumors in the pancreatic head or uncinate and local resection or distal pancreatectomy for tumors located in the pancreatic body or tail. In our case, the patient was relatively young, and the tumor was situated on the pancreatic body-tail. To preserve the function of the pancreas, central pancreatectomy was chosen. Regarding cases of pancreatic paraganglioma with metastases, Al-Jiffry et al. advocates for postoperative radiation therapy, with I131-MIBG radiation therapy being the method of choice. A large multicenter prospective study showed that high-specific-activity I-131 MIBG had a long-lasting antitumor effect in 22% of patients with advanced paraganglioma.

CONCLUSION

We reported a primary malignant paraganglioma of the pancreas, the second such case in a young patient. Draining veins may be a vital imaging sign for diagnosing pancreatic paraganglioma, and surgical resection is the primary treatment modality for these tumors. Comprehensive preoperative imaging evaluations and adequate
preoperative preparation are critical, and dual-energy imaging, such as DLCT, can provide more precise information about the lesion before surgical resection.

ACKNOWLEDGEMENTS
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Figure Legends
Figure 1 Images of the pancreatic tumor. A: On the non-contrast abdominal computed tomography (CT), white arrow indicates the strip calcification in the mass; B: Abdominal contrast-enhanced CT showed the mass was significantly enhanced, and the abundant intratumoral vessels could be seen in the tumor; C: Maximum intensity projection (MIP) of the arterial phase in a coronal view, red arrow indicates the draining vein into the portal vein; D-E: Abdominal MRI shows a low-intensity lesion on T1-weighted imaging (T1WI) and a hyper-intensity lesion on T2WI; F: MRCP shows no dilation of bile duct and pancreatic duct.

Figure 2 Spectral Imaging of the pancreatic tumor. A-D: The conventional computed tomography image (A and B), iodine density overlay (C), and Z-effective (D) images show no involvement in the gastric wall.
Figure 3 Region of interest plot of hounsfield unit attenuation. The hounsfield unit attenuation plot of the tumor (purple) and ln2 (red) and ln3 (blue) has similar slopes, that was to say that the ln2 and ln3 were most likely metastatic lymph node (confirmed by pathological examination). LN: Lymph node; HU: Hounsfield unit.

Figure 4 Histological examinations of the pancreatic tumor. A: Hematoxylin-eosin staining demonstrating the classic Zellballen pattern of paraganglioma (HE, ×100).
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<th>Ref.</th>
<th>Year</th>
<th>Age</th>
<th>Sex</th>
<th>Size (cm)</th>
<th>Location</th>
<th>CT or MRI appearance</th>
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<td>2012</td>
<td>65</td>
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Jiang et al\cite{18} 2021 41  M  4.1 × 4.2  Body  CT: A solid, No  DP  Lymph node  Present
heterogeneous soft tissue dense tumor, marked enhancement during the arterial phase and the venous phase.

Present case 2022 26  F  8.0 × 5.0 × 4.0  Body-tail  CT: A highly vascular and poor-defined mass, remarkable enhancement in arterial phase MRI, a low-intensity lesion in T1-weighted images and a high-intensity lesion in a T2-weighted images.
M: Male; F: Female; NE: Not evaluable; PD: Pancreatoduodenectomy; DP: Distal pancreatectomy; CP: Central pancreatectomy;
MRI: Magnetic resonance imaging; CT: Computed tomography.
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