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An incidentally discovered asymptomatic para-aortic paraganglioma with Peutz-Jeghers syndrome

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Peutz-Jeghers syndrome (PJS) is an autosomal dominant inherited disorder characterized by mucocutaneous melanin pigmentation and gastrointestinal (GI) tract hamartomatous polyps and an increased risk of malignancy. In addition to polyposis, previous studies have reported increased risk of GI and extra-GI malignancies in PJS patients, compared with that of the general population. The most common extraintestinal malignancies reported in previous studies are pancreatic, breast, ovarian and testicular cancers. We report the case of a 17-year-old boy who presented with generalized weakness, recurrent sharp abdominal pain and melena, had exploratory laparotomy and ileal resection for ileo-ileal intussusception. Pigmentation of the buccal mucosa was noted. An abdominal computed tomography scan (CT) revealed multiple polyps in small bowel loops. Gastroscopy revealed multiple diminutive polyps in stomach and pedunculated polyp in duodenum. Colonoscopy revealed multiple colonic polyps. Pathological examination of the polyps confirmed hamartomas with smooth muscle arborization, compatible with Peutz-Jeghers polyps. CT scan guided left para-aortic lymph node biopsy revealed the characteristic features of extra-adrenal para-aortic paraganglioma. Although cases of various GI and extra GI malignancies in PJS patients has been reported, the present case appears to be the first in literature in which the PJS syndrome was associated with asymptomatic extraaortal para-aortic paraganglioma. Patients with PJS should be treated by endoscopic or surgical resection and need whole-body screening.

Key Words: Asymptomatic para-aortic paraganglioma, intussusception, Peutz-Jeghers syndrome, screening

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CASE REPORT

A 17-year-old boy was presented to GI clinic with generalized weakness and episodic sharp abdominal pain. He had two exploratory laparotomies in 2006 and 2008 for ileo-ileal intussusception. He had no family history of GI disorders. Physical examination revealed pigmentation around the lips, face, and oral mucosa [Figure 1]. Laboratory data were
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Figure 1: Melanin pigmentation on lips and face

Figure 2: CT scan abdomen demonstrates intraluminal masses leading to intussusception (thick arrows) and a large necrotic lymph node in left para-aortic location (thin arrow)

unremarkable except for microcytic, hypochromic anemia with hemoglobin of 11.4 gm/dl. An abdominal computed tomography scan (CT) was performed revealing multiple intraluminal mass lesions, the largest one being 3.2 × 2.8 cm in small bowel loops and few of them were resulting in intussusception; however, there was no bowel obstruction noted. A large necrotic lymph node which measured 2.5 × 2.4 cm was identified in left para-aortic location [Figure 2]. CT scan-guided left para-aortic lymph node biopsy was performed and histological examination revealed a neoplastic mass composed of large polygonal cells arranged in compact aggregates with abundant finely granular eosinophilic to clear cytoplasm with distinct cell boundaries. Nuclei were round to oval and appearing mild to moderate pleomorphic and hyperchromatic. These cell nests were characteristic of paraganglioma [Figure 3]. Special glycogen stains (PAS with diastase) exhibited patchy glycogen. Immunohistochemically, tumor cells were positive for S-100 protein, chromogranin A, synaptophysin, vimentin and inhibin. Morphological and immunohistochemical features were consistent of extra-adrenal paraganglioma. His 24-hour urinary vanillyl mandelic acid (VMA) levels of 13.0 mg/24 hour (Normal: <13.6 mg/24 hour) was border line elevated. Whole body scintigraphy with 131-I labeled meta-iodobenzylguanidine (MIBG) revealed focal uptake at the level of the D12/L1 on left to mid line corresponding to the neoplastic growth noted on CT.

Gastroscopy revealed multiple diminutive gastric polyps and pedunculated duodenal polyp 1.5 cm in size, removed with snare. Colonoscopy revealed multiple pedunculated polyps about 2-3 cm in size, 10 polyps were removed endoscopically with polypectomy snare [Figure 4]. Histopathological examination revealed the characteristic PJS polyp consisting of a branching framework of connective tissue and smooth muscle lined by normal intestinal epithelium, rich in goblet cells; elongated and convoluted glands and an arborizing pattern of growth [Figure 5].

Exploratory laparotomy and enteroscopy were performed and multiple, multilobulated, broad-based polyps were excised from proximal and distal jejunum. Para-aortic circumscribed, hard 4 × 3 cm mass with feeding vessel was excised.

As a result, a final pathological diagnosis of “extra-adrenal paraganglioma” with PJS was made. The patient’s postoperative period was uneventful, and the symptoms of recurrent abdominal pain resolved. He is presently under regular and close follow-up.

DISCUSSION

Patients with PJS have an increased risk of malignancies that are not confined to the GI tract. Giardiello et al., first reported an increased risk of GI and non-GI cancers in patients with PJS.[6] The most common extraintestinal malignancies reported in previous studies are pancreatic, breast, ovarian and testicular cancers. Previous studies showed 15-fold higher incidence of cancer than the general population and 93% lifetime risk of developing any cancer among the patients suffering with PJS.[6-9]

Extra-adrenal paraganglioma is not part of the known spectrum of PJS, and thus the finding of paraganglioma and PJS may be coincidental and seems to be very rare.

Paragangliomas are rare extra-adrenal chromaffin tumors that arise from neuroectodermal cells of the autonomous nervous system and the mean age at diagnosis is reported to be about...
Paragangliomas may occur in syndromic or sporadic forms. The former is associated with well-known syndromes of multiple endocrine neoplasia syndrome (MEN): Types 2A and 2B, Von Hippel-Lindau disease, neurofibromatosis type 1, familial paraganglioma-phaeochromocytoma syndromes. The recognized genes associated with paraganglioma are: the von-Hippel Lindau tumor suppressor gene, the RET protooncogene leading to multiple endocrine neoplasia type 2, the neurofibromatosis type 1 gene, and the three genes encoding subunits B, C, and D of mitochondrial succinate dehydrogenase. The association of paragangliomas with PJS could present a new line of investigation for the ongoing genetic studies in an effort to better understand the pathogenesis of this syndrome and its associations with non-GI malignancies.

Our patient differs from those previously reported with respect to the presence of a more rare nonfunctional extra-adrenal paraganglioma. The diagnosis and management of extra-adrenal paraganglioma in a patient with PJS does not differ from the standard methods. Once the diagnosis of paraganglioma is made, every attempt should be made to perform a complete surgical resection.

Screening and surveillance programs for PJS are mandatory both for prevention of GI complications related to the polyps and for early detection of relevant malignancies. Surveillance colonoscopy, video capsule endoscopy and upper GI endoscopy should be repeated every 3 years in patients with significant polyps detected on baseline examination. However, there is no evidence for screening of other malignancies in PJS.

CONCLUSION

In conclusion, the reported case of extra-adrenal paraganglioma in a PJS patient underlines the importance of surveillance and screening programs for GI and non-GI malignancies in these patients.

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