Large operable Ewing sarcoma of the pancreas: Report of a case and review of the literature

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Abstract: Extra osseous ewing sarcoma is rare which mainly arise in paravertebral soft tissue and associated with CNS and ewing sarcoma of abdominal viscera is very rare. Only 14 to 18 cases of EES/pPNET arising in the pancreas have been reported so far in literature. We report a case 22 year boy presented with 15 days history of pain and lump abdomen. CT shows 18 *12cm mass in head of pancreas with central necrosis. CT scan showed no signs of metastatic tumor spread. Whipple procedure done for this patient, diagnosis of ewing sarcoma was confirmed by histology and immunohistochemistry. IHC tumor was positive foe neuron specific enolase, KI 67 and MIC-2 and negative for pancytoceratin and progesterone. After surgery all margin was negative 8 LN was removed which was negative for metastasis. Patient has taken chemotherapy (6 cycle of VAC-vincristine adriamycine cyclophosfamide) and doing well after 6 month of operation. CT after 3 month of surgery was normal there was no evidence of disease. so any pancreatic mass which grows rapidly and patient present with short history we should think about PNET/ES and consider it as a differential diagnosis.

Keywords: Ewing’s sarcoma, Extraosseous Ewing’s, sarcoma, Primitive neuroectodermal tumor, PNET, Pancreas

INTRODUCTION

Ewing sarcoma is after the name of James Ewing who first describe the disease as a "diffuse endothelioma of bone. Ewing sarcoma cover a huge spectrum of tumor which shows small hyperchromatic nucli withpale cytoplasm arranged in broad sheets with minimal intercellular marerial these tumor are positive for nural marker.typcally all tumor of this family shere a same genetic abnormality namly a t(11;22) translocation.

Usually ewing sarcoma is disease of bone children n young adult affected mainly, these tumor mainly compose of small round cell having hyperchromatic nucli scant pale cytoplasm. ewing sarcoma mainly metastasise through hematogenous route to lung and other bones occasionally it may arise from soft tissue (Extraosseous Ewing’s Sarcoma [4].

Two theory is given to explane its origin first ewing sarcoma arise from a primitive cell of nural crest(embryologic tissue) derived second it arise from resident cell called mesenchymal stem cells that have a capability to transform to other type of tissue become. These tumor not only shere same kind of histopathological feature under microscope but also have same kind of genetic abnormality called translocationPathologists have long known that Ewing sarcoma looks very similar to an even rarer soft tissue tumor called primitive neuroectodermal tumor (PNET). By the early 1980’s, ES and PNET were found to not only have similar features when examined under a microscope, but in greater than 95% of cases they also had an identical genetic abnormality called a translocation15. Subsequently, these two type of tumors have been grouped into a family of cancers entitled Ewing’s Sarcoma Family of Tumor (ESFT), all of which share this translocation.

In ESFT, the translocation is between chromosomes 11 and 22 and is referred to as t(11;22). The gene on chromosome 22 encodes the Ewing sarcoma gene (EWS) [16]. The gene from, named FLI1 present on chromosome 11, is responsible for turning other genes on and off. This new fused gene, called EWS/FLI, encodes an altered fusion protein that regulates other genes that can give rise to cancers when inappropriately expressed [2].

Tefft gave a series of 5 patients with a small round blue cell tumor that histologically resembled ES but arose from the paravertebral soft tissues. Since then, Extraosseous ewing sarcoma has been recognized as a distinct disease entity. that affects young adults aged 10
to 30 years, with equal sex distribution. This is more aggressive than osseous ewing it is curable, with the best prognosis seen in patients younger than 16 years of age[17].

**CASE REPORT**

A 22 year old boy presented in surgery OPD with a short history of dull aching pain abdomen and rapidly enlarging lump abdomen. On examination hard lump was present in right lumber right hypochondrium and right side of umbilical region move slightly with respiration. Patient had mild jaundice CT shows 18 *12 cm mass in head of pancreas with internal necrosis and heterogenous enhancemen and body and tail was normal. No LN seen fat plane with surrounding structure was normal spared.

EUS gave same information EUS guided FNAC taken which came to be a undifferentiated small round cell malignant tumor. Routine blood investigation was normal except bilirubin was mildly raised. Serum amylase and lipase was normal. Chest x-ray was normal. Surgery was planned and classical whipple procedure done. Per-op finding was, large solid tumor arising from head of pancreas around 20*15 cm size extending right side up to lateral abdominal wall superiorly up to porta inferiorly below umbilicus and on left side up to 6cm left to mid line. Tumor was compresing the duodenum pushing transverse colon inferiorly. There was no evidence of metastasis. Grossly, the tumor was well-circumscribed, capsulated, measured 20*15 cm, and weighed 1200g. Cut section revealed a grey tan hemorrhagica solid and cystic tumor with areas of necrosis.

**Fig-1: Large mass arising from head of pancreas. Duodenum is compressed over tumor stomach is displaced upward**
Fig-2: Large tumor displacing the SMV to the left side

Fig-3: after resection the retroperitoneal tissue shows portal vein and junction of SMV and splenic vein, right kidney is displaced inferriorly.

On histopathological examination

Macroscopically
Tumor of 15*12 cm size (largest PNET in pancreas ever reported) in head of pancreas cut surface is grayish white.

Microscopically
Section from mass reveal a cellular malignant neoplasm composed of small oval to round cell arranged in a dissociated manner nuclei show fine chromatin, occasional mitotic figure is seen, pancreatic tissue seen in one area. All (pancreatic,intestinal,gastric and CBD) resected margin is negative of tumor all LN negative of tumor metastasis(0/8), GB shows feature of chronic cholecystitis

Fig-4 & 5: Sheets of small round cells with enlarged nuclei, fine stippled chromatin, and moderately clear to amphophilic cytoplasm (H&E stain)

On IHC examination
IHC showed tumor was positive for neuron specific enolase, KI 67 and MIC-2 and negative for pancytokeratin and diagnosis was made ewing sarcoma.
Post operative course was uneventful and patient was discharged on postoperative day 6. Patient took adjuvant chemotherapy after 3 months of surgery. CT is repeated which was normal.

**DISCUSSION**

Fourteen to eighteen cases of PNET of pancreas has been reported till now [5-14]. The patients ranged in age from 6 to 25 years (mean 18 years). 60% of the patients are male. Mainly, the patients presented with jaundice and/or abdominal pain. All of the tumors were located in the head of the pancreas, and they ranged in size from 3.5 to 9.0 cm. Light microscopy revealed the typical morphologic features of PNETs. By immunohistochemistry the neoplastic cells in cases expressed O13 (CD99, p30/32MIC2) [3].

Out of 16 cases till now reported most of tumor presented with pain and jaundice [8]. Pain is most consistent symptom. [11]. 4 presented with pain and mass in abdomen. Upper GI bleeding and anemia, 2 very young girl age 2 and 6 presented with pubic hair, breast and vaginal bleeding on investigation there is increase estrogen level.

10 of the 16 tumors were located in the head of the pancreas. Tumor size varies from 3 to 15 cm in largest diameter but in our case this is the largest tumor which is reported is of 15*12 cm size (CT 18*12cm).

CT scan and MRI used to be the investigation of choice in tumor arising from pancreas in our case we also do EUS and FNAC was taken through this which was came to be undifferentiated round cell malignant neoplasm but for PNET/ES image guided core biopsy is more useful for preoperative diagnosis by HPE and IHC study. It may arise any part of pancreas but mostly been arising from head. Out of 16, 12 tumor was present in head of pancreas and periampillary lesion, 3 in body and tail of pancreas and 1 in uncinate process. Three patient presented with metastasis into the liver at initial presentation and in one tumor was infiltrated into transverse colon. Surgeries were planed according to size and site in pancreas. On microscopy, all cases exhibited same pattern, the sheets or lobules of small round tumor cells with round to oval nuclei and scant cytoplasm characteristic of the EFT. on Immunohistochemical staining all shows CD99 positive. Some proportion of the tumors stained positively for vimentin, cytokeratin and NSE. Focal Homer Wright rosettes were described in two cases. Molecular techniques such as FISH and polymerase chain reaction (PCR) demonstrated abnormalities of the EWSR1 gene in seven cases, and the t(11;22) was detected by conventional karyotyping in Movahadi 7 cases series 1 patient died of operative complication, 1 died of disease after 4 year, 1 alive with recurrence at 27 month, 1 alive with no e/o disease at 33 month and 1 who underwent biopsy and adjuvant chemotherapy alive at 43 month. Most of patient who timely underwent curative resection and taken adjuvant chemotherapy and they do well 3 to 4 year of follow up. (further followup is not available). and who did not taken adjuvant chemotherapy survival is short and recurrence is high in that cases. Overall there is no data to comment on recurrence and survival and prognosis of pancreatic PNET/ES due to lack of good long follow-up and only because few cases have been reported.

We can give an opinion that any mass arising in young patient in pancreas which is rapidly enlarging we should think of PNET/ES of pancreas as differential diagnosis. Curative resection and adjuvant chemotherapy should be the management and active follow-up for recurrence.

**REFERENCES**

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