Pancreatic Extraskeletal Ewing Sarcoma/Primitive Neuroectodermal Tumor: Case Report of a Rare Pathology

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Abstract
Ewing sarcoma, characterized by an 11:22 translocation, commonly occurs in bone and soft tissue sites. We present a patient with a history of both chest wall desmoid tumor and a subsequent adenosquamous pancreatic cancer treated with distal pancreatectomy, adjuvant chemotherapy, and radiation. Two years later the patient developed lymphadenopathy, which was biopsied and sent for the tissue of origin testing. Results revealed a sarcoma with 90% certainty, primitive neuroectodermal tumor subtype. The original pancreatic tumor was sent for next-generation sequencing, which revealed an EWSR1/FLI1 fusion as the sole genetic mutation suggestive of an underlying Ewing sarcoma. Additionally, the same fusion gene was detected on the supraclavicular lymph node biopsy. Our case affirms a change in diagnosis that was not suspected clinically with a resultant change in therapy. Though the pancreas is an unusual site of extraskeletal Ewing sarcoma (EES), emerging technologies such as next-generation sequencing can aid in management.

Keywords
Peripheral primitive neuroectodermal tumor; Ewing sarcoma; Pancreatic neoplasm

Introduction
The most common primary tumor of the pancreas is adenocarcinoma, which comprises about 85% of all cases with primary mesenchymal tumors of the pancreas being comparatively rare. Of these, Ewing sarcoma/Primitive Neuroectodermal Tumors (PNETs) are an infrequent occurrence with few such case reports existing in the medical literature. Our case discusses an unexpected diagnosis of a morphologically atypical EES found in the unusual location of the pancreas.

Case
We present a case of a 58-year old female who was undergoing surveillance for a symptomatic desmoid tumor of the chest wall previously treated with sorafenib who was incidentally found to have a 1.9 cm distal pancreatic mass. Initial biopsy suggested a tumor with neuroendocrine features. The patient subsequently underwent distal pancreatectomy with negative margins and lymph nodes. Surprisingly, the tumor was morphologically and immunophenotypically most compatible with an adenosquamous tumor evidenced by medium-sized cells growing in sheets and nests intermixed with atypical glands (Figure 1A and B), diffuse and strong labeling for squamous markers CK5/6 (Figure 1C) and P63 (not shown here). This unique, but not entirely classic histology and immunohistochemical analysis with broad keratin positivity led to a final diagnosis of a stage IIA, T3 N0 adenosquamous cell cancer of the pancreas. In addition to extrapancreatic tumor extension, perineural and lymphovascular invasion were also confirmed.

The patient received adjuvant chemotherapy with gemcitabine and 5-fluorouracil and radiation. On routine follow up two years later new supravacuicular lymphadenopathy was noted, which was restaged with PET CT showing FDG avid enlarged aortocaval, preaortic, and left axillary lymph nodes and a paraspinal soft tissue mass. These findings were suspected to be a recurrence of pancreatic cancer. The patient subsequently underwent ultrasound-guided left supravacuicular lymph node biopsy with pathology initially suggestive of a pure squamous component. Based on the unusual pattern of recurrence and the morphology that was different than the primary, Tissue of Origin (Cancer Type ID, Molecular Cancer Classifier, BioTheranostics) testing was sent, which surprisingly reported a sarcoma with 90% certainty, primitive neuroectodermal tumor subtype.

The patient was started on gemcitabine and nab-paclitaxel initially as this was suspected to be recurrent adenosquamous cell cancer of the pancreas despite the tissue of origin testing results, and additional testing on the original pancreatic tumor was sent for next-generation gene sequencing. This revealed an EWSR1/FLI1 fusion as the sole genetic mutation suggestive of an underlying Ewing sarcoma. Additionally, the same fusion gene without additional genetic mutations was detected on the supravacuicular lymph node biopsy confirming this as recurrence of her original pancreatic tumor.

Our patient was started on doxorubicin, vincristine, cyclophosphamide, and mesna with pegfilgrastim support, alternating with etoposide and ifosfamide. She completed 7 cycles and her most recent PET CT shows no evidence of metabolically active disease.

Discussion
Ewing sarcoma was first characterized to have the 11:22 translocation in 1993 [1]. Subsequently, Ewing sarcoma and Ewing-like tumors have been further characterized but novel translocations are still being identified [2-5]. Importantly, there is no agreement regarding which tumors are Ewing sarcoma, Ewing family or Ewing-like tumors. For clinical trials, exclusion criteria are increasingly being based on translocation type as it is recognized that some translocations portend a poorer prognosis such as CIC/DUX4 translocation [6-8]. Historical terms for these tumors include Primitive Neuroectodermal Tumors (PNET) and Askin tumors of the chest wall. While the tissue of origin is thought to be neuroendocrine, tumors can appear in extraskeletal sites, more often with increasing age.

Histology of these tumors reveals primitive round to oval tumor cells often with glycogen aggregates in the cytoplasm. CD99 is expressed in almost all cases of Ewing sarcoma in a characteristic...
membranous fashion, though not specific; membranous labeling by CD99 was detected in our case (Figure 1D). Vimentin stains most tumor cells and neural markers such as Neuron-Specific Enolase (NSE) are frequently expressed [9]. FLI1 is a sensitive marker for ES but is not specific. Other malignancies including lymphoblastic lymphomas, Desmoplastic Small Round Cell Tumor (DSRCT), Merkel cell carcinoma, and synovial carcinoma can show nuclear staining for this marker [10]. Some similar cases of peripheral Primitive Neuroectodermal Tumors (PNETs) of the pancreas exist in the medical literature. One large literature review by Komforti et al, 2018, documented 32 cases of histologically confirmed pancreatic Ewing sarcoma, with patients aged 2-60 with a median age of 20.5 years without significance in gender predominance [11]. Our patient, who was 58 years old at diagnosis, is among the oldest of reported patients to be diagnosed with EES of the pancreas, as the majority of reported cases occur in the second or third decade of life. Outcomes have varied among reported cases. Of 26 patients with documented follow up, two were reported alive with disease, 17 were alive without disease, and 6 died of disease with date of expiration ranging from 3 to 48 months.

Five-year survival rates in localized Ewing sarcoma have improved in recent decades due to advances in chemotherapy intensity or timing along with refinements in local control [12-14]. In patients with localized disease, those with extraskeletal sites have a greater overall survival when compared with patients with classic skeletal tumors [15-17].

Conclusion

Our case is important as it establishes the phenotypic range of this neuroendocrine tumor that typically arises in the bones of adolescents and young adults. While this clinical behavior in terms of location of metastases was not consistent with a typical relapse of pancreatic cancer, it is admittedly also unusual for EES, which typically metastasizes to lung most often and less often to other bone and bone marrow sites. Additionally, the IHC and morphologic phenotype along with the entire pathologic workup may be informative regarding a differential diagnosis for pancreatic cancers with squamoid histomorphology and immunophenotype. Increasingly, emerging technologies such as next-generation sequencing are providing unexpected clarity to unusual diagnoses.

References

defines an aggressive subtype with co-association of STAG2 and TP53 mutations. Cancer Discov 4: 1342-1353.


