FINE NEEDLE ASPIRATION DIAGNOSIS OF PANCREATIC CARCINOMA {ACINAR CELL CARCINOMA} – CASE REPORT
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ABSTRACT: Acinar cell carcinoma of the pancreas is a rare neoplasm. Pre-operative percutaneous, ultrasound guided (USG), fine needle aspiration cytology (FNAC) of pancreatic lesions is a good modality for obtaining early tissue diagnosis, aiding in distinction of clinically/radiologically appearing benign against malignant lesions. It also helps in obtaining a diagnosis in inoperable cases where chemotherapy or radiotherapy can be the line of treatment.

KEY WORDS: US-guided, Pancreas, Acinar cell carcinoma

INTRODUCTION: Ultrasound guided Fine needle aspiration cytology [FNAC] has emerged as a primary diagnostic modality in investigations of patients with pancreatic lesions. Tao et al used this technique initially to diagnose pancreatic cancer. Most solid masses of the pancreas are primary, unresectable, malignant neoplasms that show extra pancreatic extensions. Majority are of exocrine origin and have a poor prognosis. FNAC, ultrasound-guided or percutaneous, is a sensitive [81% -98%] and highly specific [99% - 100%] modality for the diagnosis of pancreatic lesions. The acini account for 80% of the pancreatic mass. However, acinar cell carcinoma are rare, comprising no more than 1% to 2% of all pancreatic neoplasms. In an population based study in USA total of 672 patients were identified in comparison to 58,526 of pancreatic adenocarcinoma which were diagnosed. They occur at any age, the median age being 51 years with a slight male preponderance. The symptoms are non-specific. Rarely acinar cell carcinoma is functional.

CASE REPORT: We are presenting a case of a 70 year old female who complained of generalized weakness, decreased appetite and altered bowel habits. On clinical examination multiple vague abdominal nodules were felt on palpation, for which USG guided FNAC was done. Free fluid was present. Clinical diagnosis offered was a colonic carcinoma with metastasis. The case was posted for exploratory laparotomy based on the clinical diagnosis. Patient was referred to sonography and fine needle aspiration before being taken up for operation. Sonography revealed a CBD(common bile duct) block and a differential diagnosis of duct carcinoma or pancreatic carcinoma with multiple secondaries was offered.

Ultrasound guided FNAC was done from the multiple irregular abdominal nodules. Multiple [5-8] slides were prepared and stained by H&E(hematoxylin and eosin), MGG(may Grünwald Giemsa), PAP(Papanicolaou) and special stains. The remaining material in the hub of
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the LP (lumbar puncture) needle was processed for cellblock. Special stains like PAS (Periodic Acid Schiff), PAS with diastase and Mucicarmine were done.

On cytology, smears were highly cellular. Tumor cells were arranged in acinar pattern, with abundant eosinophilic granular cytoplasm. The nuclei were large, crowded and contained visible prominent nucleoli. Background showed individual tumour cells with inflammatory cells and necrotic material [Fig.1]. Cells were PAS positive with resistance to diastase. Mucicarmine stain was negative.

Cell block was stained by routine H & E. The section studied showed tumour cells arranged in solid sheets and acinar patterns. Most tumour cells had scant to moderate amount of eosinophilic and granular cytoplasm, relatively uniform and centrally located nuclei which were small and had conspicuous nucleoli [Fig.2]. Based on the cytology, cell block features and also taking the clinical history with radiological features into consideration a differential diagnosis of pancreatic acinar cell carcinoma and duct carcinoma was offered.

DISCUSSION: Even though acinar cell carcinoma of pancreas account for only 1% to 2% of pancreatic carcinoma their prognosis is poor, with a 10% 5 year survival 5,8.

The closest differential diagnosis to acinar cell carcinoma is pancreatic endocrine tumours [PET], esp. Islet cell tumours. The other differential diagnosis to acinar cell carcinoma are pancreatoblastoma and solid pseudo papillary tumours of pancreas [SPT]1,4,7,9

Cytological features go against SPT and pancreatoblastoma esp. the latter which occurs in childhood4,10. Mucin stains like mucicarmine are positive in duct carcinoma. They are negative in acinar cell carcinoma as it was found in the present case4. In acinar cell carcinoma there is immunoreactivity for Trypsin, Lipase and less commonly for Chymotrypsin and Amylase. A minor endocrine component identifiable by chromogranin is present in one third to one half of the cases6,10, i.e. 30% to 40% of cases show focal positivity in acinar cell carcinoma8, while islet cell tumours show diffuse positivity with chromogranin 8. Immunocytochemistry using chromogranin was done on the cell block.

The present case showed focal positivity on cell block section, for chromogranin.

We could not send the case for electron microscopy. Thus cytomorphological features with immunocytochemistry helped to confirm the diagnosis of pancreatic acinar cell carcinoma9,11. The patient was not operated upon and was referred to a cancer centre for radiation therapy.

Thus preoperative percutaneous USG FNAC of pancreatic lesions is a good modality for obtaining early tissue diagnosis aiding in distinction of clinically and radiologically appearing benign v/s malignant lesions. A spectrum of pancreatic lesions can be diagnosed by the technique and it carries a sensitivity of 81% and specificity of 100% 1,12. Also in cases where disease is unresectable, chemotherapy, radiotherapy, or a combination of these modalities may be used to increase overall quality of life as it was noted in our case13.

CONCLUSION: USG guided FNAC of pancreas is a safe, efficacious method to obtain material from pancreatic tumours. It also helps to give diagnosis by cytology along with ancillary methods like cell block and immunocytochemistry. It is a useful diagnostic aid in surgically inoperable cases.

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FIG1 FNAC showing tumour cells arranged in acinar pattern(H &E, 400)

FIG2 chromogranin positivity seen on cell block confirming the diagnosis of acinar cell carcinoma pancreas.(400x)