Diffuse replacement of pancreatic parenchyma by intraductal papillary mucinous neoplasm

A 63-year-old insulin-dependent diabetic woman presented to our multidisciplinary group for evaluation and management of multifocal cystic lesions of the pancreas found incidentally by computed tomography during an evaluation of diverticulitis. Initial assessment included endoscopic ultrasound (EUS) with fine-needle aspiration (FNA) by curved linear array echoendoscope. Near-total replacement of the pancreatic parenchyma by innumerable cysts of various size and character was demonstrated, involving all regions of the pancreas (Video 1). Several cysts exhibited features that gave cause for concern, including size well over 30 mm, thickened walls, and thickened septations, as well as round, mobile solid structures suggestive of mucin balls. Moreover, the main pancreatic duct appeared to communicate with the cysts in the regions of the head and neck (Video 1). The minimally spared parenchyma was heterogeneous without lobularity. The major vessels of the abdomen were unremarkable and without abnormality, nor was there evidence of other regional disease or lymphadenopathy. FNA was performed to recover fluid contents from a larger cystic component, and a second pass was performed to confirm the mobility and benign nature of a suspected mucin ball.

Video 1
Endoscopic ultrasound demonstrating near-complete replacement of the entire pancreatic parenchyma.

Fig. 1 a–e Endoscopic images demonstrating near-complete replacement of the pancreatic parenchyma: a uncinate process and head, b head, c neck, d body, e tail.
Fluid analysis returned with relatively high levels of both amylase (26,900 U/L) and carcinoembryonic antigen (CEA; 589 ng/mL), and associated cytopathology was paucicellular without suspicious cells, although a thick mucus component was noted. Serum tumor marker evaluation included a cancer antigen (CA) 19–9 concentration of 1739 U/mL and a CEA concentration of 3.4 ng/mL. Interval pancreatic protocol computed tomography supported the EUS findings of near-complete replacement by multiple cystic lesions without evidence of regional or metastatic disease (Fig. 3). The finding of multiple intraductal mucinous cysts is consistent with a diagnosis of diffuse side-branch intraductal papillary mucinous neoplasm (IPMN). The surgical procedure was uncomplicated, and the patient recovered and was discharged with follow-up care as anticipated.

This case presents a unique, previously unreported demonstration of near-complete pancreatic replacement by diffuse side-branch IPMN diagnosed by EUS and confirmed with surgical excision. Accurate differentiation between various etiologies of cystic pancreatic lesions presents challenges even in the most typical of scenarios [1]. By recognizing sometimes subtle distinguishing characteristics, however, the differential diagnosis may frequently be narrowed to one solitary etiology. Here, EUS alone predicted the diagnosis in part by allowing fluid analysis. The study demonstrated a multifocal distribution of thickly septated cysts clearly communicating with the main pancreatic duct, which when aspirated showed a viscous clear fluid rich in CEA and amylase in an elderly patient. While not definitively exclusive, this pattern differs from those expected for mucinous cystic neoplasms (body/tail of pancreas, solitary, multiloculated, ovarian-like stroma, without ductal communication, low amylase), serous cystadenomas (head of the pancreas, solitary, microcystic/sponge-like, central fibrous scar sometimes calcified, without ductal communication, low amylase and CEA), pseudocysts (single, unilocular, patient with history of pancreatitis, low CEA), solid–pseudopapillary neoplasms (body/tail, young female, solid and cystic, without ductal communication, low amylase and CEA), lymphoepithelial cysts (elderly male, cystic, associated with keratinous debris, without ductal communication, low amylase and CEA) and cystic neuroendocrine tumors (nonseptated, without ductal communication, low amylase and CEA) [2]. Given the diffuse nature of the cystic lesions, von Hippel–Lindau disease was also considered; however, this syndrome is associated with serous cystadenomas and/or neuroendocrine lesions, not mucinous neoplasms, as demonstrat-

**Fig. 2** a Well-defined, round structure within a larger cyst; on fine-needle aspiration it was found to be mobile b.

**Fig. 3** a, b Contrast-enhanced computed tomography confirming near complete replacement by cystic lesions: a transverse and b coronal views.
ed by the cytopathology obtained on FNA [3]. Current consensus guidelines for management of IPMNs dictate consideration of total pancreatectomy to address the diffuse lesions, which are suspicious for progression to malignancy due to their size and worrisome features [4].