Combination of Right Nephrectomy and Total Pancreaticoduodenectomy for von Hippel-Lindau Disease

Nikolaos Arkadopoulos1, Konstantinos Karapanos1, Vaia Stafyla1, Anneza Yiallourou1, Andreas Koureas2, Agathi Kondi-Pafiti3, Vassilios Smyrniotis1

Departments of 1Surgery (Second), 2Radiology, and 3Pathology, Aretaieion Hospital, Athens University School of Medicine. Athens, Greece

ABSTRACT
Context Von Hippel-Lindau disease is an inherited syndrome of multorgan neoplasia caused by a germline mutation in the von Hippel-Lindau gene and can include central nervous system tumors, renal cell carcinomas and benign pancreatic cystic tumors. Case report We report the case of a 56-year-old patient who had a past history of cerebellar hemangioblastoma and presented with abdominal pain. Imaging revealed renal tumors and multiple pancreatic tumors which caused duodenal and pancreatic duct compression. The patient was treated with a combination of radical right nephrectomy, total pancreaticoduodenectomy and splenectomy. Pathology identified a multifocal unilateral clear cell renal carcinoma which interestingly coexisted with multiple large pancreatic serous microcystic adenomas with infiltration of the fibrous capsule. Conclusion In past cases of von Hippel-Lindau disease, pancreatic adenomas with malignant transformation have not been reported. In our case, the infiltration of the fibrous capsule by parenchymal cells may indicate malignant transformation.

INTRODUCTION
Von Hippel-Lindau disease is an inherited syndrome which usually presents in the second decade of life and causes neoplastic lesions in multiple organs. Renal cell carcinoma is the main malignant tumor and the leading cause of death while pancreatic lesions are usually benign. We present the case of a 56-year-old female with von Hippel-Lindau disease who was diagnosed with renal cell carcinoma and synchronous multiple pancreatic lesions causing duodenal obstruction and chronic pancreatitis due to pancreatic duct compression. The patient underwent a right nephrectomy, total pancreaticoduodenectomy and splenectomy. Histology revealed clear renal cell carcinoma and serous pancreatic cystadenomas, with focal invasion of the fibrous capsule, a finding which is interesting in the light of the benign nature of these lesions.
pancreatic lesions. Endoscopic ultrasound-guided FNA biopsy of the lesion on the pancreatic head showed severe chronic pancreatitis. Pancreatic neuroendocrine hormones and urinary metanephrine excretion were within normal range. With a clinical diagnosis of von Hippel-Lindau disease, the patient underwent genetic evaluation which was positive for mutation R161P in exon 3 of the von Hippel-Lindau gene, in heterozygosis (direct sequencing).

Surgical exploration was performed through a bilateral subcostal incision. A large solid mass on the superior pole of the right kidney and another smaller similar lesion on the inferior pole were confirmed. The pancreatic parenchyma was multilobular and of woody texture, indicative of chronic pancreatitis. A large hard mass in the pancreatic head was severely compressing the second part of the duodenum close to the ampulla, without invasion of the surrounding organs. After intraoperative evaluation of these findings, we proceeded with an en-bloc total pancreatico-duodenectomy and splenectomy, followed by a radical right nephrectomy. The patient had an uneventful postoperative course and was discharged on the 7th postoperative day. Histological examination showed two well-differentiated clear cell renal adenocarcinomas in the upper and lower poles of the right kidney. The pancreas measured 15x5x4.5 cm, had a woody texture and contained one hard, microcystic mass measuring 5x5x4 cm in the head, and one multilobulated cystic lesion in the body and tail, measuring 5x2x2 cm. Both lesions proved to be serous microcystic adenomas with cellular and architectural atypia. The adenoma of the pancreatic head harbored a small focus of microscopic infiltration of its fibrous capsule (Figure 3). The rest of the pancreatic parenchyma exhibited severe changes characteristic of chronic pancreatitis, as the result of significant compression of the pancreatic duct by the two large serous microcystic adenomas.

DISCUSSION

Von Hippel-Lindau disease is an inherited syndrome caused by deletions or mutations in a tumor suppressor gene (VHL gene) mapped to human chromosome 3p25 [1]. Affected individuals can develop central nervous system lesions including cerebellar, spinal cord, brainstem, nerve root and supratentorial hemangioblastomas as well as retinal hemangioblastomas and endolymphatic sac tumors. The main visceral manifestations of the disorder include renal cysts and carcinomas, pheochromocytomas, pancreatic cysts, neuroendocrine tumors and serous cystadenomas as well as epididymal and broad ligament cystadenomas. Its clinical manifestations present in the second decade of life and its penetrance reaches 90% by the age of 65 years [2]. Renal cell carcinoma is the main malignant tumor in von Hippel-Lindau disease and is usually the leading cause of death. Its prevalence is estimated between 24 and 45%. The overall incidence of renal lesions (including renal cysts) is 60% [2, 3, 4], with the mean age at presentation being 39 years. Renal lesions are often multiple, multifocal and bilateral. They usually remain asymptomatic for long intervals and
only in the more advanced cases do they become clinically apparent. Treatment recommendations depend on tumor size. Nephron-sparing surgery is recommended for carcinomas having a maximum diameter of 3 cm [4]. Nephron or renal-sparing resection is designed to reduce the risk of metastasis while preserving kidney function. Large carcinomas (greater than 3 cm), especially when multifocal disease is confirmed, carry an increased risk of metastases. In rare cases, when the kidney cannot be preserved, a total nephrectomy is the only option. In our case, a large unilateral multifocal renal cell carcinoma, located in both poles of the right kidney, obliged us to perform a radical right nephrectomy.

The prevalence of pancreatic lesions of von Hippel-Lindau is 35-70%, reaching up to 93% of affected individuals in some studies. The most common lesions in the pancreas are cystic (17-56%), with a notable incidence of serous microcystic adenomas [5, 6, 7]. Nevertheless, solid tumors may also develop, such as neuroendocrine tumors (8-17%) and less frequently adenocarcinomas. The mean age at presentation for pancreatic lesions of von Hippel-Lindau is 35-37 years. They are rarely the presenting feature of the disease and usually remain asymptomatic. However, it has been reported that serous microcystic adenomas can cause local compressive effects, particularly at the level of the papilla of Vater [8]. Although these same lesions are thought to be predominantly benign, there have been reports of malignant microcystic adenomas [9]. In our case, a large solid mass in the head of the pancreas, with suspicious characteristics on preoperative imaging, was compressing the second part of the duodenum as well as the main pancreatic duct, causing severe symptoms. The decision to perform a total pancreatectomy was mainly based on the patient’s severe symptomatology and the intraoperative findings of chronic pancreatitis. However, most von Hippel-Lindau cases can be treated with more conservative pancreatic surgery.

In conclusion, we have herein presented an unusual case of von Hippel-Lindau disease with the unique combination of renal cancer and large pancreatic serous microcystic adenomas, one of which was found to focally invade the fibrous capsule. The pancreatic lesions were the cause of coexisting severe chronic pancreatitis, due to compression of the pancreatic duct. In such cases, the proposed combination of nephrectomy and pancreatectomy in a single operation appears to be a safe and oncologically comprehensive approach.

Conflict of interest The authors have no potential conflict of interest

References