Hepatocellular Carcinoma with Congenital Absence of the Portal Vein in a Child

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Abstract

Congenital absence of the portal vein (CAPV) is such a rare malformation of the splanchnic venous system. Also it can be a reason for hepatic tumors, for example adenoma, hepatoblastoma or hepatocellular carcinoma (HCC). The cooccurrence of CAPV and HCC in adult patients have been reported in the literature. It is the second case of a pediatric patient whom has also CAPV and HCC.

Introduction

Congenital absence of the portal vein (CAPV) and its intra-hepatic branches is a malformation of the splanchnic venous system described by John Abernethy in 1973 [1]. Almost thirty cases have been reported to date [2,3]. CAPV is generally seen in children and associated with multiple anomalies for example skeletal abnormalities, cardiac failure and hepatic tumours [4]. The co-occurrence of CAPV and hepatocellular carcinoma (HCC) has been described in three adults and in only one child [5,6]. We present a pediatric patient with concomitant occurrence of HCC and CAPV.

Case Report

A 5 years old girl was admitted to pediatrics department of another hospital general with pollakiuria and loss of appetite since 2 years. Abdominal ultrasoundography showed an iso/hyperchoic liver mass about 35 mm involving segment 7 which has a hypoechoic halo and peripheric vascularisation. Then the patient was referred to pediatric oncology department of our hospital for further investigation. She has a history of term birth with normal birth weight and she has no significant medical history. There was no history of malignancy in her family.

Her weight and height values were in normal percentiles. The pathological findings were bilateral submandibular lymphadenopathies about 1.5 cm and 2/6 systolic murmur at mesocardiac region. Liver and spleen were not palpable. Liver function tests were as follows; AST 45 U/L (N<35), ALT 29 U/l (N<40), Alkaline phosphatase 201 U/L (N<110), LDH: 2198 U/L (N<220) and total bilirubin: 0.76 mg/dl (<1.10). The ammonia level was normal. Initial a-fetoprotein was 14 g/L (N), so we had ruled out metabolic disorders.

Computed tomography revealed two solid lesions, larger one about 1.5 cm and 2/6 systolic murmur at mesocardiac region. Liver and spleen were not palpable. Liver function tests were as follows; AST 45 U/L (N<35), ALT 29 U/l (N<40), Alkaline phosphatase 201 U/L (N<110), LDH: 2198 U/L (N<220) and total bilirubin: 0.76 mg/dl (<1.10). The ammonia level was normal. Initial a-fetoprotein was 14 g/L (N), so we had ruled out metabolic disorders.

According to The European staging system, considers only the pretreatment extent of disease, (PRETEXT, the PRETreatment EXTent of disease scoring system), our patient was Stage II because tumor-free and also base structure of the parenchyma was protected.

Computed tomography revealed two solid lesions, larger one localized in the segment 8 of the liver. The lesion was extending from the dome of the liver to the portal hilum. Right hepatic vein was compressed and displaced to posteriorly. Central hypervascular nodular enhancement was observed in the arterial phase of CT imaging. The smaller lesion was located in the segment 6. It was isodense with the liver parenchyma in the arterial phase and hypodense on delayed venous phase images. The CT examination also revealed an absence of the main portal vein (Figure 1). Superior mesenteric vein was coursing left of the midline at the level of the head of pancreas and after joining with the splenic vein at the level of the pancreatic tail they both were draining into the hemiazygous vein. No pathological uptake was observed on PET scans. Echocardiography was normal.

Ultrasound-guided fine-needle aspiration biopsy was performed from the segment 8 lesion. Reticulin network loss and sinusoidal CD34 expression were preserved within the tumor that supports the malignancies but staining pattern with β-catenin does not support malignancies. The pathology report of the biopsy material was “atypical hepatocytotic nodule” and surgical excision of the mass had been suggested. Thus, the patient underwent surgery and right hepatectomy was performed. Surgeon noted that extrahepatic portal vein and its right and left branches were absent. Standart right hepatectomy was done without any difficulty.

Microscopic examination demonstrated “Multifocal hepatocellular carcinoma, grade II” noting surgical margins and the liver capsule was tumor-free and also base structure of the parenchyma was protected. According to The European staging system, considers only the pretreatment extent of disease, (PRETEXT, the PRETreatment EXTent of disease scoring system), our patient was Stage II because tumor

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Figure 1: Abdominal contrast-enhanced computed tomography showing the absence of main portal vein.

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involves two adjoining quadrants (segments 6 and 8). According to the post-surgical North American staging system, our patient is stage I which means no metastases and tumor completely resected [7].

Two months after surgery, patient received two cycles of chemotherapy POG-8697 regimen which including Cisplatin (100 mg/m² iv-infused-Day 1), Vincristin (1.5 mg/m² iv-inf Day 3,10,17) and 5-flourourasil (600 mg/m² iv-Day 3). According to a POG study 3 years evidence free survey is 91% for stage I and II patients which treated with POG-8697 regimen [8].

Two months later, in order to evaluate the response to treatment, computed tomography was performed and no recurrent or residual lesion was detected. The patient is doing well during 8 months follow-up.

Discussion

Malignant liver tumors are seen rarely that's about<1% of all pediatric malignancies [7]. Contrary to adults, in whom the predominant malignancy is hepatocellular carcinoma, hepatoblastoma accounts for 70% in children. Hepatocellular carcinoma is the most frequent hepatic malignancy in adolescents. Generally, hepatic viral infection like HBV, HCV or cirrhosis is related with hepatocellular carcinoma and it can take decades for malignancy to develop, because of this it is seen in very young children rarely. CAPV is another reason for HCC in adults. There are only a few pediatric cases who has HCC associated with CAPV in the literature [5,6].

CAPV is an uncommon anomaly in which the splanchnic venous flow is diverted away from the liver and drains into the systemic circulation. It is fewy symptomatic and other congenital abnormalities which are cardiovascular like dextrocardia, ventricular oratrial septal defects, patent foramen ovale and ductus arteriosus; skeletal such as hemivertebrae, fifth finger anomalies, ocuolaocularulvertebral dysplasia; gastrointestinale for example biliary atresia, liver tumors and urinary like cystic renal dysplasia, hippospadia, vesico-urethral reflux can be associated [9-12]. Liver tumors have been described in the presence of distracted portal venous flow and they can be benign (focal nodular hyperplasia, adenoma) or malignant lesions (hepatoblastoma, HCC) [13,14].

If we consider how CAPV cause liver tumor; hepatotrophic factors such as pancreatic hormones carried by portal system may be the answer. They may initiate regeneration of hepatic cells [15]. With the absence of portal hepatic perfusion results in change of hepatic cell structure [16]. Absence of portal venous flow may be compensated by a stronger arterial flow then it leads to the development of intrahepatic nodular lesions[17].

In our case, we could not find any infectious, metabolic, or genetical reason that causes HCC except CAPV. Interestingly, our patient is 5 years old and there is not long time to develop HCC. That made us think CAPV may be affected the case since fetal life and/or type of CAPV also may be important for developing HCC.

Conclusion

Since our patient is one of the rare cases which reported in the literature, a relationship between portal vein agenesis and HCC needs to be explained with more cases with multicenter studies. In children, HCC is usually associated with metabolic diseases, usually after progression of the disease to cirrhosis. There also is an increased prevalence of HCC in children in areas where there is high hepatitis B virus exposure [18]. Our patient did not have an underlying cirrhotic liver and was negative for hepatitis B virus and she received hepatitis-B vaccine in infancy. Here, we would like to stress the importance of hepatitits B vaccination for help protect yourself from hepatitis B infection especially in developing countries.

On the other hand follow-up protocols and preventive procedures should be established for these children. Early diagnosis of recurrent disease of our patient is important since vascular anomaly of portal vein may be still affecting. Cadaveric liver transplantation can be done if the patient has recurrent disease during follow-up.

References