CASE REPORT

Inferior Vena Cava Hypoplasia Associated with Deep Vein Thrombosis – Case Presentation

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Introduction
Inferior Vena Cava (IVC) malformations such as absence of infrahepatic IVC or IVC hypoplasia are rare, being recorded in approximately 0.5% of the general population [1]. This condition is usually asymptomatic, the diseases being diagnosed incidentally, associated with other congenital anomalies such as situs inversus, congenital heart diseases, polysplenia and asplenia [2]. During physical exertion, the collaterals that are formed in the evolution of the disease are unable to withstand the increased blood flow, thereby generating venous stasis and clotting. Since during embryogenesis right metanephrons drain into the IVC, congenital absence of the latter could affect right kidney development. Some authors have suggested this condition as KILT syndrome (Kidney and IVC abnormalities with Leg thromboses) [3].

Case report
We present the case of a male patient, aged 58 years, who presented with intense pain in the lower limbs, with the onset 24h before presentation. The patient presented generalized edema, cyanosis and functional impotence. From his personal history, we mention that the patient is suffering from essential hypertension stage II with high cardiovascular risk and non-insulin necessitating type II diabetes with renal and vascular complications. The physical examination revealed an influenced general condition, warm skin, pain, swelling, cyanosis of the lower limbs, generalized subdiafragmatic edema and bilateral basal crackles stasis.

Laboratory tests on admission were within normal limits. Urinalysis did not show any pathologic changes. ECG on admission showed no significant changes. The EcoDoppler examination of the lower limbs revealed a dilated inferior vena cava, with intraluminal hypoechogenic material, bilateral dilated external iliac veins, with intraluminal hypoechogenic content. The common femoral veins were dilated, with the lumen occupied with blood clots. The right femoral vein had a complete thrombosis on the whole length and an acute deep vein thrombosis completely occluding distal inferior vena cava was noted. Abdominal ultrasound revealed hepatic steatosis, chronic pancreatitis, hepatomegaly and kidney tumor formation on both kidneys. Also, at admission, an abdominopelvic AngioCT was performed, indicating hypoplasia of the inferior vena cava with azygos vein dilatation (fig. 1), perirenal venous dilatation (fig. 2), bilateral (on the right side pronounced) and infrarenal extensive DVT (of interest for the VCI, VIC internal and external, bilateral VFC); delayed bilateral renal function, bilateral renal cortical cysts; enlarged bilateral renal sinus (hydronephrosis). Concluding all clinical and paraclinical examinations, the patient was diagnosed with phlegmasia cerulean dolens and hypoplasia of the inferior vena cava.

Thrombolytic therapy was initiated (streptokinase for 72h), associated with anticoagulants (Heparin), with a significant amelioration in the clinical status. The circumference of the leg/thigh reduced and we also recorded a significant pain relief and reduction of inflammation signs on the lower limbs. After 8 days of treatment, AngioCT...
examination was repeated and found an arterial abdomi-
nal pelvic system and lower limb arterial system without
detectable lesions (fig. 3), renal excretion present bilater-
ally, peritesticular dilated varicose veins associating bilat-
eral hydrocele (fig. 4) and deep venous system undergoing
reperfusion with thrombotic debris at the level of inferior
vena cava, common iliac vein confluence, left common
iliac vein, bilateral internal iliac veins, right iliac vein, all
associated with minimum peritoneal fluid collection.

Discussions
Patients with inferior vena cava hypoplasia are prone to
develop deep vein thrombosis due to the venous stasis of
the lower limbs [4]. In the case described, the patient had
iliofemoral DVT of the lower limbs, which could be re-
lated to the hypoplasia of the vena cava inferior, revealed
by imagistic evaluation. The presence of spontaneous,
recurrent and sometimes bilateral proximal lower limb
DVT must call the physician’s attention to the possibility
of an inferior vena cava anomaly [5]. MRI and contrast
CT angiography are especially useful in such cases. How-
ever, DVT as a paraneoplastic manifestation or as a result
of a hypercoagulable state should also be excluded. The
most appropriate treatment in such cases is anticoagu-
lation for at least six months. The possibility of recurrence
is high when the anticoagulation treatment is discontin-

Fig. 1. Hypoplasia of the inferior vena cava

Fig. 2. Perirenal venous dilatation

Fig. 3. Lower limb arterial system

Fig. 4. Peritesticular dilated varicose veins
ued before this period. Patients with IVC hypoplasia can present with different types of clinical pictures. Some can be asymptomatic and hypoplasia of the IVC is found as an incidental finding, while others can present with venous thrombosis and its consequences as in our case.

**Conclusion**

Vena cava inferior atresia complicated with a deep vein thrombosis has a good prognosis under treatment with anticoagulants and thrombolytics.

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**References**