Case history

A 53 year old male vegetarian, not diabetic or hypertensive and did not have a history of cardiovascular, hepatic or kidney disease, presented to us with vague pain and worsening swelling of both lower limbs since 1 week. Physical examination was unremarkable except for bilateral tender pitting edema of feet, legs and thigh. Investigations showed Hb 116g/L, WBC count 8.9 X 10^9/L, platelet 301 X 10^9/L, ESR 72mm/hr, blood urea nitrogen 21mmol/L, serum creatinine 132 micromol/L, Urine albumin nil. Saturation was 98% in room air. Chest X ray was unremarkable. Venous doppler study of lower limbs unmasked DVT involving bilateral saphenofemoral, superficial popliteal and popliteal veins with a high total plasma homocysteine of 29.76umol/L (5.46-16.20umol/L). CT Venogram showed chronic thrombosis of the IVC and bilateral renal veins with acute thrombosis in bilateral common iliac, external/internal iliac, bilateral common femoral and superficial femoral veins. Colonoscopy showed multiple colonic diverticulae. 2-D ECHO showed myxomatous AML suggestive of mitral valve prolapsed (MVP).

Keywords: polycystic kidneys, deep vein thrombosis, homocysteine, colonic diverticulae, mitral valve prolapse

Abstract

A 53 year old male presented with vague pain and worsening swelling of both lower limbs since one week. Venous doppler study of lower limbs unmasked DVT involving bilateral saphenofemoral, superficial popliteal and popliteal veins with a high total plasma homocysteine of 29.76umol/L (5.46-16.20umol/L). CT Venogram showed chronic thrombosis of the IVC and bilateral renal veins with acute thrombosis in bilateral common iliac, external/internal iliac, bilateral common femoral and superficial femoral veins. Colonoscopy showed multiple colonic diverticulae. 2-D ECHO showed myxomatous AML suggestive of mitral valve prolapsed (MVP).

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Figure 1 CT Venogram showed hypoplastic IVC as depicted by thinned out IVC with chronic thrombosis(white arrows) with acute thrombosis in bilateral common femoral/superficial femoral veins depicted by distended veins when compared to the arteries(black arrows).

CT Abdomen with contrast showed right kidney 10X6.1cm with >10cysts, Left kidney 9.6X5.1cm with 3-4 cysts, the largest cyst on either side measuring about 6cm (Figure 2). Tc-99m Isotope renogram scan revealed total GFR 65ml/min (left kidney 52ml/min, right kidney 13ml/min).
Figure 2 CT Abdomen with contrast shows Right kidney 10X6.1cm with >10 cysts, Left kidney 9.6X5.1cm with 3-4 cysts, the largest cyst on either side measuring about 6cm with Tc-99m Isotope renogram revealed total GFR 65ml/min(left kidney 52ml/min, right kidney 13ml/min).

Prostate malignancy was ruled out. Plasma CEA and CA19-9 were normal, stool occult blood negative, upper GI endoscopy showed gastric ulcer with duodenal erosions with biopsy from lesions ruling out malignancy, colonoscopy showed multiple colonic diverticula (Figure 3).

Figure 3 Colonoscopy showing colonic diverticuli(arrow).

2-D ECHO showed good biventricular function, EF-62%, type 1 diastolic dysfunction, myxomatous AML suggestive of mitral valve prolapse (MVP) (Figure 4).

He was treated with Inj. Enoxaparin 60mg s/c BID followed by oral warfarin to maintain the PTINR between 2 and 3. Patient also received additional B6 and folic acid for hyperhomocystinaemia.

Figure 4 2-D ECHO showing myxomatous AML (arrow) suggestive of mitral valve prolapse (MVP).

Discussion

Venous thrombosis and hypercoagulability is a common complication of malignancy and hypoplasia of the IVC, described as an incidental finding, should be taken into account when otherwise unexplained DVT develops in young people. Hyperhomocysteinemia is a risk factor for recurrent venous thrombosis in patients between 20 and 70 years of age. Our patient had a hypoplastic IVC as depicted by thinned out IVC with mild to moderate hyperhomocystinaemia explaining the probable cause for venous thrombosis. However an extensive workup to detect malignancy was negative.
ADPKD is associated with MVP in up to 25%. There are rare case reports of thrombosis in the inferior vena cava (IVC) due to enlarged cysts in autosomal dominant polycystic kidney disease. Our patient showed number of cysts to meet the required criteria for ADPKD, with Stage II CKD, MVP, multiple colonic diverticulae and IVC thrombosis, had no positive family history, with only elder brother having late onset minimal change disease. Genetic workup for ADPKD is done. Literature review did not show association of IVC hypoplasia with cystic kidney disease.

Acknowledgments

None

Conflicts of interest

Authors declare there is no conflict of interest.

References