Abnormalities of gut vessels in Turner's syndrome

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Summary

We describe a 57-year-old patient with Turner's syndrome, iron deficiency anaemia and intestinal vascular abnormalities. Colonoscopy revealed 2 widely dilated, tortuous veins in the terminal ileum and several smaller ectatic veins and haemangioma-like malformations throughout the colon. Laparotomy for herniotomy showed only minimal vascular abnormalities of the serosal surface. Patients with Turner's syndrome and anaemia should be checked for these lesions by endoscopy, and conversely, in patients with such lesions, Turner's syndrome should be considered.

KEY WORDS: Turner's syndrome, anaemia, gut vessel abnormalities.

Introduction

In 1938, Turner published his classical paper describing a congenital disorder associated with gonadal dysgenesis, webbed neck and cubitus valgus. Further abnormalities associated with the same syndrome have been described since then (Haddad and Wilkins, 1959). An association of Turner's syndrome with gastrointestinal haemorrhage due to haemangiomas, was described by Lisser et al. (1947). Since then, 23 cases have been reported, most of them having been discovered at laparotomy. Diagnosis was made by endoscopy in a few cases only (Rutlin et al., 1981; Rosen, Sirota and Marinoff, 1967). We describe a patient with Turner's syndrome and gastrointestinal bleeding probably due to vascular abnormalities, discovered by endoscopy.

Case report

A 57-year-old unmarried kitchen aid was admitted to the hospital in October 1981 with anaemia. In 1972 appendicectomy had been performed. In 1974, at the age of 50, diagnosis of Turner's syndrome with hypogonadism, primary amenorrhoea and a typical dysmorphic habitus was made. Sex chromatin was completely absent in leucocytes, in the mucosa of the mouth and in the roots of the hair. Hormonal substitution was not given. In 1978, she had a rightsided inguinal hernia repair which recurred a few weeks later. She presented in July 1981 with a 3 month history of fatigue and was found to be anaemic. The haemoglobin was 9.0 g/dl, and the reticulocyte count was 0.8%. The white cell count and platelets were normal.

No stainable iron was found in the bone marrow. Stool examination for occult blood was positive 3 times. On gastroscopy, several spots of intestinal metaplasia were found in the antrum and proven by histological examination. Colonoscopy showed 2 very tortuous, varix-like veins in the terminal ileum, several centimeters in length with a diameter of about 7 mm (Fig. 1). There was no active bleeding during endoscopy. Throughout the colon several similar abnormalities were found, although much smaller in size. Furthermore, some blue strawberry-like prominent abnormalities with a maximum diameter of 5 mm were observed. A selective small bowel series revealed no abnormality; varix-like veins could not be detected.

The patient was discharged on iron replacement therapy. One month later the haemoglobin had risen to 11.9 g/dl. At operation to repair the hernia, the serosal surface of the terminal ileum and colon showed only minor vascular abnormalities, some very thin but tortuous vessels near the mesentery. Tortuous veins shone through the intestinal wall in the ileum (Fig. 2).

Discussion

The intestinal vascular abnormalities seen in this patient seem to be sufficiently characteristic to raise
Clinical reports

Fig. 1. Endoscopic view of tortuous varix-like vein in the terminal ileum (diameter about 7 mm) in a patient with Turner's syndrome. (Endoscope: Olympus CF IBW, videocamera: ITC Ikegami. Tsushinki Co. Ltd., Japan.)

Fig. 2. Tortuous veins shining through the intestinal wall of the ileum at laparotomy in the same patient. The serosal layer is normal.
the possibility present in female patients of Turner’s syndrome (Rosen et al., 1967). Similar vascular abnormalities have not been described in men. Other causes of ectatic veins, such as portal hypertension, were excluded. To our knowledge, only 23 such cases have been published to date. Haddad and Wilkins (1959) examined 56 patients with Turner’s syndrome and found a prevalence of vascular lesions of about 5 to 8%. Ectatic veins or haemangiomas have been described, predominantly on the serosal side (Haddad and Wilkins, 1959; Rosen et al., 1967). In most patients, only one type of abnormality has been described. Rosen et al. (1967) first suggested that all these varieties could be the expression of the same vascular anomaly. In fact, we found different vascular abnormalities in our patient, dilated veins and venules at different sites in the colon and the ileum with extremely gross varicose veins in the ileum (so-called ‘bag of worms’), telangiectasia and several haemangioma-like malformations. In contrast to other reports (Haddad and Wilkins, 1959; Rosen et al., 1967), the lesions were only on the mucosal side. The endoscopic finding of these vascular abnormalities in the terminal ileum has not yet been reported.

Routine X-ray examination of the gastrointestinal tract, including selective small bowel series, is not adequate to demonstrate these vascular malformations (Frame et al., 1977), as in our case. Angiography was not performed in the present case, and, with the size of the vessels seen, some abnormality might have been expected.

Bleeding seems to be intermittent, and intestinal haemorrhage apparently tends to decrease in severity and frequency with age (Frame et al., 1977). A regression of these vascular abnormalities with age was shown in 2 patients by Frame et al. (1977) and Scott (1978). Electrocoagulation or the injection of sclerosing agents have not to our knowledge been tried. In rare cases surgery may be needed. Considering the benign course of the lesions we believe that patients should be treated with iron replacement and blood transfusions if necessary.

References


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