INTRODUCTION
Mirror image transposition of abdominal and thoracic viscera is termed situs inversus. Total situs inversus affects both chest and abdomen while partial situs inversus affects one region only or both regions in an incomplete manner (2). Situs inversus is detectable in 1/4000 to 1/20,000 live births (2). Patients are susceptible to life threatening defects of the heart and abdomen, including cardiac chamber defects, transposition of the great vessels, intestinal malrotation, bowel and biliary atresias and splenic anomalies (3, 4).

Obstruction of the duodenum is rare in situs inversus. Nawaz et al traced only 18 cases in the medical literature prior to 2004 (5). That study pointed to annular pancreas or a duodenal diaphragm causing obstruction in the majority of patients. Approximately 40% of situs inversus cases subjected to laparotomy have a preduodenal portal vein (PDPV) but this vein almost never causes obstruction (3, 6).

We herein report a case of duodenal obstruction in situs inversus caused by a PDPV.

CASE REPORT
A three-day-old full term female baby presented with bilious vomiting from birth. The infant weighed three kilograms, was moderately dehydrated and had a non-distended abdomen. Babygram X-rays done on admission showed a reverse double bubble gas pattern (gastric bubble on the right and duodenal bubble on the left) and dextrocardia (Fig. 1). These features, along with limited intestinal gas, suggested incomplete duodenal obstruction and total situs inversus. Echocardiography confirmed dextrocardia in a heart that was otherwise normal. After correction of dehydration and hypotension (sodium-124 mmol/L), laparotomy was performed through a left upper abdominal transverse incision. The duodenum, biliary tree and caecum were found to be situated in the left abdomen and the descending and sigmoid colon in the right.

The duodenal C-loop was incompletely formed and tethered by adhesive bands (Ladd’s bands) in keeping with intestinal malrotation. The junction between the first and second part of the duodenum was compressed by a PDPV, causing proximal dilatation and distal collapse (Fig. 2). The adhesive duodenal bands were divided and an appendectomy done. Duodenotomy revealed no luminal obstruction. The
obstructing PDPV was bypassed by constructing a side-to-side duodeno-duodenostomy between dilated and collapsed duodenum.

Postoperatively, the baby had tonic clonic seizures presumed to be due to hyponatraemia (125 mmol/L) and hypocalcaemia (1.50 mmol/L). This was controlled by phenobarbitone and normal saline infusion. Duodenal paresis caused a delay in the commencement of feeds until day 12 post-surgery but subsequent recovery was uneventful leading to discharge on day 18.

DISCUSSION

Aristotle first detected situs inversus in animals and considered it a visitation from the gods (2–4). Fabricius uncovered the condition in cadavers in 1600 (1, 2) but the clinical significance of situs inversus was not grasped until the advent of X-ray imaging which made diagnosis in humans easy and reliable (1, 2). Clinical syndromes with a definite link to situs inversus have been defined. These include asplenia (predominance of right sided structures) and polysplenia (predominance of left sided structures) syndromes (Ive-mark’s syndrome) (3), multiple organ malrotation (MOMS) syndrome (levoversion of liver, dextroversion of stomach and normal large and small intestines) (7) and Kartegener’s syndrome (total situs inversus, bronchiectasis and chronic sinusitis) (3). All but the last are frequently associated with life-threatening intra-abdominal and cardio-pulmonary anomalies (3, 4, 7).

A PDPV rarely causes duodenal obstruction (6, 8, 9). It derives from a caudal anastomotic vein connecting the two vitelline veins at the 5 mm embryo stage. Non-regression of this vein leads to a ventrally routed portal vein (8). Surgeons should anticipate a PDPV in situs inversus and ascertain its presence by ultrasonography, computed tomography (CT) or magnetic resonance imaging (MRI) (3, 6). Steps should be taken to avoid injury to this vein because of possible life-threatening consequences (8). Duodenal obstruction by a PDPV is best treated by an intestinal bypass procedure such a duodeno-duodenostomy (6, 8, 9).

The main causes of duodenal obstruction in situs inversus were found to be annular pancreas, duodenal webs, adhesive bands and midgut volvulus (3). This case had both a PDPV and duodenal adhesive bands linked to intestinal malrotation. The PDPV was diagnosed as the source of duodenal obstruction because the vessel crossed the duodenum at an anatomical transition point between proximally dilated and distally collapsed duodenum. Furthermore, no intraluminal obstruction was detected after duodenotomy and passage of a catheter into the distal duodenum (3, 10).

Where intestinal malrotation is found at laparotomy, a modified Ladd’s procedure (lysis of duodenal adhesions and appendectomy) should be performed (3, 5, 10) to reduce the risk of future adhesive obstruction and acute appendicitis. The latter is difficult to diagnose in situs inversus. One study reported false pain projection to the right in 33% of such patients with appendicitis (2). This false pain projection along with confounding symptomatology leads to misdiagnosis and wrongly placed incisions (2, 3).

The potential for cardiac pathology is so great in situs inversus and the consequences so unpredictable that all surgical cases should be subjected to echocardiography prior to anaesthesia (5).

REFERENCES


