Giant chest wall arteriovenous malformation with spinal cord compression

Sir,
Congenital arteriovenous malformations (AVMs) may occur in any organ. Skin AVMs are common, but AVMs of the chest wall or extracranial space are uncommon. We report a patient with an extensive AVM that involved skin, chest wall, and extracranial space. Definitive treatment of such an AVM is difficult.

A 23-year-old man presented with progressive lower extremity weakness. A large cutaneous ‘haemangiomatous’ of the right chest posterior wall measuring 22 cm in greatest diameter had been present since birth (Figure 1). There was no history of ulceration, bleeding, or rapid increase in size. Underlying soft tissues of the chest wall were involved. The AVM was easily compressible and it readily refilled. Neurological examination showed generalised weakness of both lower extremities. The right plantar response was extensor; left plantar response was flexor. Laboratory investigations were normal. Chest radiograph showed deformity of the right 6th through 10th ribs. Chest computed tomography (CT) showed a large vascular mass involving the skin, chest wall, pleura, and posterior mediastinum of the right chest (Figure 2). CT myelogram showed extension of this mass into the extradural space at the T-6 level. The neural foramen was enlarged. At midthoracic laminectomy a very vascular extracranial mass was removed, confirmed to be a benign arteriovenous malformation. The patient made a complete neurological recovery. Because of the extent of the AVM, definitive surgical excision of the chest wall and mediastinal components was not recommended. Embolisation will be reserved for recurrence of symptoms.

Chest wall1 and extradural2 AVMs are uncommon. Extradural AVMs are usually located in the mid-thoracic region. They may extend into the mediastinum and paraspinal areas in a ‘dumb bell’ fashion but involve the posterior mediastinum and paraspinal space (AVMs) of the chest wall is rare.3,4 Our patient’s AVM was much more extensive than any previously reported case.

Extradural AVMs with cord compression require neurosurgical intervention.1 Extradural AVMs are usually asymptomatic but have a high risk of complication, and this was the case in our patient. The patient’s AVM could not have been resected without sacrificing approximately 50% of his chest wall. However, the patient ultimately went on to have safe resection and was discharged home without the need for postoperative haemorrhage. Careful operative haemostasis is required to prevent worsening of cord compression by extradural haematoma.

Indications for treatment of chest wall AVMs include disfigurement, bleeding, and congestive heart failure. Small chest wall AVMs can be cured by excision5 but complete resection of extra-thoracic chest wall AVMs is problematic. Our patient’s AVM could not have been resected without sacrificing approximately 50% of his chest wall. Intraoperative haemorrhage can be massive in these circumstances. In addition, radical surgical extirpation does not always result in permanent AVM control; recurrence may still occur. Arterial embolisation is the treatment of choice for very large AVMs that are symptomatic.6 Embolisation provides control of large chest wall AVMs but permanent cure is infrequent. Since embolisation is primarily palliative in cases of large chest wall AVMs, it should probably be reserved for symptomatic patients.

Figure 2 CT of the chest and spine shows a large vascular mass involving chest wall, mediastinum, and extradural space.

Figure 1 Chest wall arteriovenous malformation

Benign spontaneous pneumoperitoneum in an elderly patient treated medically with recovery

Sir
Spontaneous pneumoperitoneum signals perforation of a hollow viscus in over 90% of cases1 and patients usually undergo emergency surgery. However, in the absence of sepsis or perforation and in the presence of co-existing illnesses often seen in the elderly, laparotomy and an anesthetic might sometimes actually compromise the chance of survival. It has been suggested that these patients can often be treated without laparotomy provided that abdominal tap is negative for sepsis or inflammatory fluid.2

A frail 93-year-old with no medical history apart from her arthritis for which she took ibuprofen, was admitted with a short history of abdominal pain and diarrhoea. On examination, the abdomen was distended but there was little tenderness and she was apyrexial. Abdominal X-ray showed massive pneumoperitoneum. The surgeons recommended conservative measures. She was treated with intravenous fluids and antibiotics. Her pneumoperitoneum resolved and she was eventually discharged home without the need for surgery.

Pneumoperitoneum without perforation is a distinct entity3 which does not require urgent surgical intervention, but may cause diagnostic confusion and sometimes causes unnecessary laparotomy. It has been described as secondary to ruptured pneumatoceles cystic in the intestines,4 pneumomediastinum, pneumothorax, and sometimes following endoscopic procedures such as percutaneous endoscopic gastrostomy.5 Hence a careful history and exclusion of other causes may prevent a hazardous surgery in elderly patients with this condition.

A HUSSAIN
JGC COX
Department of Medicine, Wansbeck General Hospital, Northumberland, UK

Correspondence to JD Urschel, Division of Thoracic Surgery, Roswell Park Cancer Institute, Elm and Carlton Streets, Buffalo, NY 14263-0001, USA


References


Pneumoperitoneum: causes

• pneumatoceles cystic in the intestines
• pneumomediastinum
• pneumothorax
• following percutaneous endoscopic gastrostomy
• peritoneal dialysis
• following laparotomy

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