Submucosal esophageal hematoma following sclerotherapy: A rare complication

RAM SINGH, YK CHAWLA, UPJEEET KAUR, NEELAM MALIK, JB DILAWARI

ABSTRACT: A cirrhotic patient who developed an intramural hematoma of the esophagus as a complication of esophageal variceal sclerotherapy is reported. The hematoma resolved spontaneously within two weeks without any residual complications such as esophageal stricture. Can J Gastroenterol 1990;4(1):23-25

Key Words: Hematoma, Portal hypertension, Sclerotherapy

Hématome oesophagien sous-muqueux consécutif à une sclérothérapie: Une complication rare

RESUME: Un patient cirrhotique a développé un hématome intramural de l'oesophage, complication d'une sclérothérapie administrée dans le traitement des varices oesophagiennes. L'hématome s'est résolu spontanément en l'espace de deux semaines sans aucune complication résiduelle — sténose oesophagienne, par exemple.

Endoscopic sclerotherapy is now considered an important therapeutic modality in the management of variceal hemorrhages. There is now enough data to show that endoscopic sclerotherapy significantly reduces rebleeding risk (1-3); however, its effect on the survival of patients with cirrhosis is still controversial (2-4). Various complications of endoscopic sclerotherapy have been described in the literature (5,6). This report deals with an extremely unusual complication.

CASE PRESENTATION
A 55-year-old female presented in September 1986 with massive hematemesis and melena, which were diagnosed to be variceal in origin on endoscopy. She had normal liver function tests; however, a biopsy of the liver showed cirrhosis. After initial control of the bleed using a Sengstaken-Blakemore tube, the patient was started on endoscopic sclerotherapy (EST). EST was performed with 1% sodium tetradecyl sulphate at three-week intervals using an Olympus fibrescope. The varices decreased in size from grade III to grade II after six sessions.

During the seventh session of sclerotherapy, one grade II varix was injected with 1 mL of the sclerosant. The patient did not complain of pain during or immediately after the procedure. Over the next 12 h, however, she developed a progressive dysphagia to both solids and liquids including saliva, along with severe retrosternal pain.

On examination there was no hepatosplenomegaly and the lungs were clear. The patient had a normal coagulogram which was done soon after the complication occurred.
Figure 1) Left Barium swallow done the day after the complication, showing a filling defect which gives the esophagus the appearance of having a double lumen. Right Barium swallow showing a normal looking esophagus after two weeks of conservative therapy.

An esophagogram with gastro conray (Conray 280; May & Baker, India) (Figure 1) showed an apparent double lumen of the esophagus. There was no extramural leakage of the dye. Chest radiodiogram was normal. Upper gastrointestinal endoscopy revealed narrowing of the esophageal lumen 24 cm from the incisors with a dusky bluish nodular elevation completely occluding the lumen of the esophagus, suggestive of a submucosal hematoma. A computed tomography scan of the thorax revealed that the hematoma was confined to the submucosa of the esophagus, producing a double lumen effect (Figure 2).

The patient was kept nil by mouth and started on intravenous fluids. After 48 h she showed some improvement, being able to swallow liquids. After two weeks she could swallow solids without discomfort. The patient remained afebrile over the two week symptomatic period. A repeat esophagogram after two weeks revealed a normal esophagus (Figure 1) while endoscopy showed an area of hyperemia at the site of hematoma extending from 24 to 32 cm, without any narrowing of the lumen of the esophagus. A repeat computed tomography scan of the thorax confirmed the disappearance of the hematoma (Figure 2).

**DISCUSSION**

This case demonstrates a most unusual complication of sclerotherapy. The sudden and rapidly progressive dysphagia which developed within hours of EST, and the finding of a large dusky bluish submucosal lesion obstructing the lumen of the esophagus made hematoma a likely diagnosis, which was supported by a computed tomography scan.

Awareness of the occurrence of this hematoma as a complication of sclero-
therapy will aid in the recognition of this condition after sclerotherapy, presenting with a sudden onset of severe retrosternal pain and dysphagia. Similar symptoms have been previously reported (6, 7).

When an intravariceal injection is attempted using a free hand technique, some extravasation of the sclerosant into the surrounding tissues is inevitable (8).

Necrosis and hemorrhage into submucosal tissue from the submucosal vessels may have caused the hematoma following sclerotherapy.

Management of such lesions should be conservative, as is evident from this case study. Spontaneous resolution has been known to occur without any residual defects such as stricture or dysmotility.

Intramural hematomas of the esophagus have also been described in the literature following protracted vomiting in the absence of preceding instrumentation (9, 10). Such hematomas usually occur in the distal esophagus following a Mallory-Weiss laceration. Patients with impaired coagulation may also develop a hematoma without having vomited beforehand.

REFERENCES