Gastroenterology

SIMON P.L. TRAVIS

DPhil, FRCP Consultant Gastroenterologist John Radcliffe Hospital Oxford, UK

TARIQ AHMAD

DPhil, MRCP Specialist Registrar John Radcliffe Hospital Oxford, UK

JANE COLLIER

MD, MRCP Consultant Hepatologist John Radcliffe Hospital Oxford, UK

A. HILLARY STEINHART

MD, FRCP(C) Head, Combined Division of Gastroenterology Mount Sinai Hospital Toronto, Canada

THIRD EDITION



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Preface to the third edition

The field of gastroenterology continues its rapid pace of growth in the understanding of disease processes and the discovery of new diagnostic and treatment strategies. The success of the first and second editions of Pocket Consultant in Gastroenterology published in 1991 and 1998, in concert with significant advances in gastroenterology and hepatology over the past 5-6 years, has led to this third edition. The book's enduring appeal is a tribute to the original co-authors, Dr George Misiewicz and Rodney Taylor. Dr. Simon Travis is the leading force in this very useful book, and he has enlisted the fresh perspective of new authors, including Drs. Jane Collier, Tariq Ahmad and Hillary Steinhart. This third edition very effectively captures advances in new knowledge regarding the pathophysiology of gastrointestinal and hepatobiliary diseases and the management of patients with these disorders that has developed since the second edition. The authors continue to use the user-friendly format of short, focused paragraphs, and bulleted lists, supplemented by tables and figures where appropriate. The authors have struck the balance of remaining concise, yet being thorough in their discussions of all major disease entities. The whole story is provided in an economical format that makes this handbook so useful to busy clinicians seeking information in the fast pace of contemporary practice. This remarkable little book should be of value to primary care physicians, trainees in internal medicine and gastroenterology, as well as busy gastroenterology consultants, by providing a rapid and efficient way of refreshing themselves on specific topics. The font and layout of this book makes it particularly easy to seek out specific information. This third edition of Pocket Consultant in Gastroenterology is a goldmine of current information distilled into a terse and didactic style, making it indispensable in the clinic and office of the generalist and specialist gastroenterologist.

> Emmet B Keeffe, MD Professor of Medicine Stanford University School of Medicine President of the American Gastroenterology Association 2003-4

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> Simon Travis Tariq Ahmad Jane Collier Hillary Steinhart

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1.1 Swallowed foreign body

Toddlers, the mentally disturbed and the elderly most commonly swallow foreign bodies. If no history is available, look for excessive salivation, regurgitation, choking or distress. Objects impact in the pharynx, lower end of the oesophagus, or pylorus. Pain or fever suggests perforation. Once through the pylorus, spontaneous passage is the rule, but perforation can occur in the ileocaecal region.

All cases

- · Look in the mouth
- If the object is impacted in the fauces, call the ENT surgeon
- X-ray the chest and abdomen, but failure to visualise an object does not exclude its presence
- Look for surgical emphysema, mediastinal and subdiaphragmatic gas on X-ray
- Barium or Gastrografin examination is not indicated and may hinder endoscopy
- Address the underlying issues to avoid repeated ingestion, especially in prisoners or the mentally disturbed who may have ingested foreign bodies for individual gain

Bones, pins, glass and batteries

- Chest pain suggests perforation, and a small haematemesis may herald perforation of a major vessel. In either case, contact the thoracic surgeon urgently
- Sharp objects should be removed by an experienced endoscopist, unless they have passed the duodenum. A plastic sleeve over the endoscope helps prevent trauma during withdrawal. If possible, the endoscopist should practice snaring or grasping a similar object before starting the procedure in order to determine the best means of retrieval
- After endoscopic removal, further chest pain may indicate delayed perforation
- Batteries, especially small alkaline batteries ingested by toddlers, should be retrieved immediately if in the oesophagus. Corrosive perforation or heavy metal intoxication has been reported. Once in the small intestine, safe passage is the rule

Coins, beads and blunt objects

- Almost always pass spontaneously unless more than 5 cm long or 3 cm in diameter
- Reassure the patient or parents and advise them to check stools for 3 days
- Repeat abdominal X-ray after 36 h if there is doubt about progress. Documented arrest by X-rays for 72 h is an indication for surgical exploration

Body-packing (ingested packets of drugs)

- Smuggled packets of drugs may be swallowed, or secreted per rectum
- Intact packets can cause intestinal obstruction and if packets do not pass within 72 h, surgical removal is advisable

1.2 Complete oesophageal obstruction

- Endoscopic removal is contraindicated because of the risk of rupturing the bags
- Packets may burst spontaneously and cause life-threatening overdose
- Heroin overdose causes constricted pupils, bradypnoea or coma. Hypoglycaemia or non-cardiogenic pulmonary oedema may occur later. Give intravenous naloxone 0.8 mg rapidly, to a maximum of 2.4 mg if necessary
- Cocaine causes dilated pupils, tachycardia and agitation. Convulsions, metabolic acidosis or coma may occur. Sedate with intravenous midazolam 5–10 mg and give oral propranolol 40 mg three times daily for a few days
- Severe overdose of any narcotic is an indication for ventilation and surgical removal of the packets, to stop drug absorption
- The doctor's immediate duty is the treatment of the patient if body-packing is discovered. Once treatment has been initiated, the appropriate authorities should be notified according to local legal regulations
- Questioning of the patient must wait until the patient is fit, and be sanctioned by the most responsible physician

1.2 Complete oesophageal obstruction

Bolus obstruction causes sudden, complete dysphagia for solids and liquids, with inability to swallow saliva. Food impacted against a benign or malignant stricture is the usual cause. Occasionally the presentation is delayed for a few days in the mentally handicapped or severely debilitated. The obstruction must be relieved urgently.

Clinical features

Ask about and look for:

- Duration of symptoms preceding obstruction
- Predisposing disease (stricture, carcinoma, Schatzki's ring)
- Triggering factors (steak, toast, fibrous foods, tablets)
- · Dehydration
- Weight loss (suggests malignant obstruction)
- Supraclavicular nodes (from a carcinoma of the cardia)
- Complications (aspiration pneumonia, perforation)

Investigations

The endoscopist should be contacted as a priority.

- · Full blood count—anaemia suggests carcinoma
- · Serum electrolytes—high urea indicates dehydration
- Chest X-ray—look for a mediastinal fluid level (obstruction), absent gastric air bubble (obstruction) or right lower lobe consolidation (aspiration)
- Urgent endoscopy—must be performed by an experienced endoscopist
- A barium swallow risks aspiration and is inappropriate, unless the diagnosis is in doubt. This is *not* the same as in dysphagia without obstruction (Section 2.1, p. 51)

Management

- · Intravenous fluids
- Endoscopic removal of the obstructing bolus

1.3 Oesophageal rupture

- Endoscopic dilatation can be done immediately after disimpaction
- Carbonated drinks occasionally disimpact fibrous debris, but endoscopy is needed when a food bolus is stuck for a few hours
- Fine-bore nasogastric feeding, or nutritional supplements are needed (Section 12.2, p. 378) if dilatation is delayed after removal of the bolus. Endoscopic placement of the tube is awkward, but indicated if it cannot be inserted in the normal way (p. 413)
- Intravenous metronidazole 500 mg and cefuroxime 750 mg three times daily for 5 days, if aspiration pneumonia is present

Prevention

Simple measures decrease the risk of acute obstruction in patients with oesophageal strictures or prosthetic oesophageal tubes (pp. 59 and 65):

- · Avoid fibrous food (apples, oranges), steak and toast
- · Wear dentures if edentulous
- · Chew all solids well
- · Carbonated drinks with meals
- Avoid oral potassium supplements, salicylates and large tablets>
- A proton-pump inhibitor (PPI; e.g. omeprazole 20–40 mg, lansoprazole 30 mg or rabeprazole 20 mg daily) delays or prevents restricturing in most patients and heals associated oesophagitis. PPIs should be continued indefinitely

1.3 Oesophageal rupture

Sudden chest pain after forceful vomiting is the cardinal symptom when the distal posterior oesophageal wall tears longitudinally in spontaneous perforation (Boerhaave's syndrome). Traumatic perforation after instrumentation or chest injury is more common than spontaneous rupture.

Differential diagnosis

Perforation presents with chest pain, respiratory distress, painful swallowing or subcutaneous emphysema. Early diagnosis is crucial to survival. Failure to consider the possibility is the commonest reason for misdiagnosis.

- Myocardial infarction (ECG, cardiac enzymes)
- Dissecting aneurysm (pulses, chest X-ray, urgent CT scan)
- Perforated peptic ulcer (rigid, silent abdomen, erect chest X-ray)
- Acute pancreatitis (amylase more than fourfold elevated)
- Spontaneous pneumothorax (chest X-ray in expiration)

Investigations

Confirm the diagnosis and site of perforation.

- Chest X-ray—look for mediastinal or subdiaphragmatic gas, or a hydro/pneumothorax (Fig. 1.1)
- Gastrografin swallow—in spontaneous rupture, tears are usually large and leak contrast; after instrumental rupture, tears are often small and do not leak contrast. Upper oesophageal perforations tend to leak into the mediastinum; mid-oesophageal perfo-

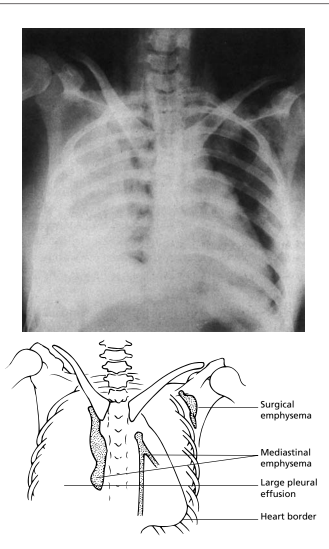


Fig. 1.1 Chest X-ray in oesophageal rupture showing consequences of oesophageal perforation. There is a large right pleural effusion, mediastinal emphysema and gross surgical emphysema in the neck and upper chest wall.

rations into the mediastinum and right pleura; and distal oesophageal perforations into the mediastinum, left pleural cavity or abdomen

Management

Oesophageal perforation is a potentially lethal condition. Conservative management is confined to highly specific situations. If the perforation has involved the pleural cavity, or has been contaminated by saliva, gastric contents or food, then surgery is mandatory.

Resuscitation

· Intravenous fluids

1.4 Caustic oesophageal injury

- Analgesia—intravenous morphine 2.5–5 mg every hour until pain is relieved, then every 4 h
- Involve thoracic surgical colleagues at an early stage

Spontaneous rupture

- · Nil by mouth
- Surgical repair and drainage is almost always needed and should occur within 24 h
- Antibiotics (intravenous metronidazole 500 mg and cefuroxime 750 mg three times daily for 5 days)
- Enteral nutrition through a jejunostomy fashioned at the time of surgery is best

Instrumental rupture

- Small tears (with minor symptoms and no leakage of contrast) may be managed conservatively in conjunction with the surgeons. Large tears that leak contrast are managed as for spontaneous rupture
- · Nil by mouth
- Nasogastric aspiration for 3 days
- · Intravenous fluids
- · Antibiotics as above
- Indications for surgery are a persistent fever, or pneumothorax after 48 h
- When perforation complicates palliative treatment of a malignant stricture, patients who
 are unfit for surgery may be managed by endoscopic insertion of a cuffed oesophageal tube

1.4 Caustic oesophageal injury

Ingestion of caustic cleaning fluids can cause progressive and devastating injury to the oesophagus and stomach. Most occur as accidents in children under 5 years. Symptoms and the appearance of the pharynx do not correlate with the extent of oesophageal or gastric injury.

Clinical features

- There may be no symptoms initially, but diagnosis is not difficult if an accurate history is obtained
- Identify the specific fluid ingested
- · Hoarseness and stridor indicate pharyngeal or laryngeal injury
- Painful dysphagia or haematemesis indicate oesophageal oedema and ulceration. This
 may occur rapidly or be delayed for several hours
- Respiratory distress and shock occur when mediastinitis develops due to oesophageal necrosis
- Acute symptoms may resolve, to be followed by dysphagia after several weeks or months as scar tissue causes an oesophageal stricture

Investigations

- Chest X-ray—look for mediastinal air or oedema
- Direct laryngoscopy and endoscopy—to record the extent of injury. This should be performed under general anaesthetic, since sedation is unsatisfactory in children and

1.5 Acute bleeding: upper gastrointestinal tract

does not allow adequate views of the larynx or hypopharynx. Circumferential burns lead to strictures

Management

Resuscitation

- · Nil by mouth
- Establish the airway by intubation or tracheostomy if respiratory symptoms or stridor are present
- · Intravenous fluids
- Do not give neutralising agents

Minor, non-circumferential burns

- · Allow liquids and then a light diet if tolerated
- · Arrange psychiatric assessment if ingestion was a suicide attempt

Deep ulcers or circumferential burns

- Remain nil by mouth for 48 h
- Monitor for chest pain or fever, indicating delayed perforation or mediastinitis. If these occur, give intravenous antibiotics, liaise with thoracic surgical colleagues and start nasojejunal feeding
- · Allow liquids and a light diet if asymptomatic after 48 h
- Omeprazole 40 mg daily, lansoprazole 30 mg daily, rabeprazole 20 mg daily or equivalent PPI for 4 weeks may reduce the rate of late stricturing, but steroids have no effect

Late complications

- Oesophagogastric strictures occur in about 25%, but almost invariably in those with deep ulcers or circumferential burns
- Review patient after 4 weeks and at 3, 6 and 12 months
- · Arrange a barium swallow if dysphagia occurs, before oesophageal dilatation
- Repeated dilatation is frequently necessary. Shortening of the oesophagus may promote gastro-oesophageal reflux, so omeprazole 40 mg daily, lansoprazole 30 mg daily or equivalent PPI is reasonable to reduce restricturing
- · Attention to nutrition is vital
- Surgery is indicated if patients cannot tolerate repeated dilatation
- · The risk of carcinoma of the oesophagus is substantially increased

1.5 Acute bleeding: upper gastrointestinal tract

The aims of management in upper gastrointestinal bleeding are to stabilise the patient, stop active bleeding and prevent recurrent bleeding. There are about 90 admissions per 100 000 adults annually in the UK, with an overall mortality of 14% unless patients are admitted to a specialised bleeding unit, where a mortality of about 5% can be expected. Most bleeds from peptic ulcers stop spontaneously and about 25% can be identified who have no risk of rebleeding and can be rapidly discharged. The task is to distinguish the 20% who rebleed in hospital and who may need surgical intervention. A standard

1.5 Acute bleeding: upper gastrointestinal tract

clinical approach is recommended for every patient, so that patients at highest risk of rebleeding and death are identified early.

Clinical approach

- Assess severity
- · Resuscitate
- Establish the site of bleeding
- · Liaise with the surgical and intensive care teams on call
- Medical intervention
- Early surgery when appropriate

Assessment

The aim is to identify patients at high risk of rebleeding and death, by clinical and endoscopic examination. The Rockall score is an independently validated risk assessment score that is simple to apply and recommended. All patients with haematemesis or melaena must be treated actively until a stable baseline has been established. There is no room for complacency.

Rockall score

The Rockall score is applied in two stages. First, there is a clinical score to be performed upon arrivals that estimates mortality:

Criterion		Score	T '.' 1 1' 1
• Age	<60 years	0	Initial mortality risk
	60-79 years	1	score (pre-endoscopy)
	> 80 years	2	0 = 0.2%
• Shock	None	0	1 = 2.4%
	Pulse & sBP > 100	1	2 = 5.6%
	sPB < 100	2	3 = 11.0%
• Comorbidity	None	0	4 = 24.6%
	Cardiac/any major	2	5 = 39.6%
	Renal/liver/malig.	3	6 = 48.9%
• Total initial score	C	(max = 7)	7 = 50.0%

• Then, after endoscopy the mortality score is updated to produce a final score:

Criterion	Score	T2 1
• Endoscopic diagnosis		Final mortality score
No lesion, or M-W tear	0	(after endoscopy) 0 = 0.0%
All other diagnoses	1	0 = 0.0% 1 = 0.0%
Malignancy of upper GI tract	2	2 = 0.2%
• Stigmata of recent haemorrhage		3 = 2.9%
None/haematin	0	4 = 5.3%
Clot, visible vessel, blood in stomach	2	5 = 10.8%
• Final score after endoscopy	(max = 11)	6 = 17.3%
		7 = 27.0%
		8 = 41.1%

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Document the following

In addition to a record of the assessment of the patient:

- Preceding symptoms (dyspepsia, vomiting, weight loss)
- · Drug and alcohol ingestion
- · Presence or absence of melaena on rectal examination
- Signs of chronic liver disease (Table 5.2, p. 137)

Causes

See Table 1.1.

Investigations and management

Resuscitation on arrival

- Ensure a patent airway
- Insert one 14- or two 18-gauge intravenous cannula
- If pulse > 100 b.p.m., give 500–1000 mL colloid (such as Haemaccel, Gelofusin, Pentaspan or pentastarch) over 30–60 min and repeat if necessary whilst waiting for blood
- Transfuse blood until haemodynamically stable in the first few hours, because initial haemoglobin is a poor indicator of the severity of the bleed. Subsequently transfuse up to haemoglobin of 10 g/dL
- Synthetic colloid or crystalloid will cause haemodilution: 1000 mL decreases the pretransfusion haemoglobin by about 10%
- Reserve group O rhesus negative blood for dire emergencies (such as continuing massive bleeding and systolic BP < 80 mmHg despite 1000 mL intravenous colloid), when the risk from hypotension exceeds that from uncrossmatched blood
- Insert a urinary catheter in patients who need a central venous line (p. 11), to monitor urine output for information regarding fluid balance
- Ensure that the patient remains nil by mouth until endoscopy
- Do not insert a nasogastric tube, because this increases the risk of haemorrhage from gastric and oesophageal lesions
- Admission to a designated specialised unit for gastrointestinal bleeding reduces mortality to 5% or less. If this is not available, consider admission of patients with a predicted mortality > 10% (initial Rockall score ≥ 3) to a critical care unit and contact surgical colleagues as soon as the patient is resuscitated

Table 1.1 Differential diagnosis of haematemesis or melaena

Common	Less common (<5%)	Rare (1%)
Duodenal ulcer (35%) Gastric ulcer (20%) Gastric erosion (6%) Mallory–Weiss tear (6%) No lesion found (20%)	Duodenitis Oesophageal varices Oesophagitis Tumours	Hereditary telangiectasia Aortoenteric fistula Haemostatic defect Pseudoxanthoma elasticum Haemobilia Pancreatitis Angiodysplasia Portal hypertensive gastropathy

1.5 Acute bleeding: upper gastrointestinal tract

Initial investigations

- Full blood count, crossmatch, coagulation studies and electrolytes
- Crossmatch 4 units of blood for patients with > 10% mortality risk (Rockall ≥ 3), but group and save alone for lower risk patients. Note that this is a practical application of the Rockall score, but it has not been validated for this purpose
- Haemodynamic status is a better guide to transfusion requirements than measured haemoglobin
- Arterial gases in those with cardiorespiratory disease
- ECG in high-risk patients
- Chest X-ray in high-risk patients (abdominal films rarely help)

Indications for a central venous line

- Signs of major haemorrhage (pulse > 100 b.p.m., systolic BP < 100 mmHg). Reasons for not inserting a central line in patients with a high (> 10%) predicted mortality (p. 9) must be carefully considered
- Rebleed during the same admission
- Inadequate peripheral venous access
- If a central venous line is needed, monitoring in a critical care area is advisable

Establish site of bleeding

- Arrange endoscopy after resuscitation, ideally within 12–24 h. Mucosal lesions and stigmata for rebleeding are otherwise missed. Ensure that the presence or absence of stigmata (p. 11) is recorded
- Indications for emergency endoscopy are continued bleeding, a rebleed in hospital, or if the patient is being considered for surgery
- Profuse haemorrhage may obscure the bleeding site. Gastric lavage to remove clots rarely alters management and can be hazardous. Repeat endoscopy after a further 12 h resuscitation is recommended. Immediate surgery should be a joint decision between surgeons and physicians
- Table 1.1 (p. 10) shows the differential diagnosis
- Interpret the endoscopy report intelligently: it should identify stigmata of recent haemorrhage (predicts risk of rebleeding), state whether there has been intervention (e.g. sclerotherapy of ulcers) and describe the position of ulcers (posterior duodenal ulcers overly a branch of the gastroduodenal artery that can rebleed vigorously)
- Stigmata of recent haemorrhage in the base of an ulcer and risk of rebleeding are:

Stigma	Risk of rebleeding
None	< 1%
Haematin (black spots)	5%
Adherent clot	30%
Visible vessel	50%
Bleeding vessel	80%

• Endoscopic intervention (below) halves, but does not abolish, the risk of rebleeding

Monitoring and discharge

- Pulse, BP, central venous pressure and urine output hourly, until stable
- · Re-examine after 4 h
- Coagulation studies if > 4 units transfused
- Daily full blood count, urea and electrolytes for patients being transfused and for 2 days after
- Keep 2 units in the blood bank for 48 h after bleeding has stopped
- Patients who do not have endoscopic stigmata of high rebleeding risk (p. 11) and who
 have not had endoscopic intervention can safely start eating and drinking immediately after endoscopy and be discharged at any time thereafter, as long as there is adequate support at home
- Patients who have had endoscopic intervention (p. 11) should be kept in hospital for 72 h after bleeding has stopped

Medical intervention

These measures are not an alternative to surgery if an operation is indicated (p. 15), but may help stop bleeding or reduce the risk of rebleeding. Endoscopic intervention is indicated for patients with a peptic ulcer and active bleeding or non-bleeding visible vessel.

- Endoscopic intervention: all techniques halve the risk of rebleeding, but depend on local expertise and may not be available. Injection of adrenaline (up to 10 mL 1: 10 000) around peptic ulcers, thermocoagulation and laser photocoagulation all have similar efficacy. Sclerosant (ethanolamine) is best avoided as necrosis and perforation have been reported. A combination of adrenaline (1: 10 000) and thrombin (1000 U/mL) may be more efficacious at preventing rebleeding
- Intravenous omeprazole or pantoprazole (20-80 mg in 250 mL saline infused over 1 h, then 8 mg/h for 72 h) is only indicated after endoscopic intervention for bleeding peptic ulcers. It halves the risk of rebleeding and surgery and reduces mortality by one-third
- Ranitidine, oral omeprazole or other acid-suppressing drugs have no place in the initial treatment of bleeding. They should be reserved for treatment once a peptic ulcer has been diagnosed
- Tranexamic acid (1 g intravenously, three times daily for 72 h) has been shown on metaanalysis to reduce rebleeding and mortality. In the absence of a previous thromboembolic event, it is a reasonable adjunct in the treatment of high-risk patients until further trials are available. It also reduces the risk of recurrent bleeding from angiodysplasia
- Determination of *Helicobacter pylori* status should be done at the time of emergency endoscopy, using a biopsy urease (*Campylobacter*-like organism, CLO) test in patients with a bleeding ulcer (p. 87). Eradication therapy for *H. pylori*-positive patients (table 3.2, p. 89) is indicated as soon as oral feeding is restarted
- Terlipressin (Glypressin) 2 mg bolus, then 2 mg every 4 h is indicated for varices (below); Glypressin may not be available in all countries
- Other drugs (octreotide, vasopressin) do not have a proven role in the management of acute non-variceal gastrointestinal bleeding. A combination of ethinyloestradiol (50 µg) and norethisterone (1 mg/day) may decrease episodes of recurrent acute bleeding from angiodysplasia (such as hereditary telangiectasia)

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Rebleeding

Rebleeding greatly increases mortality. Patients at high risk of rebleeding (based on endoscopic stigmata, p. 11) need to be recognised and the surgeons told of their admission. Patients are best admitted to a critical care area, where signs of rebleeding should be detected early. Signs of rebleeding are:

- Rise in pulse rate (a sensitive and early sign)
- Fall in central venous pressure
- Decrease in hourly urine output
- · Haematemesis or continued melaena
- Looking at the patient (pallor, pulse, postural pressure drop and poor peripheral circulation) is as important as looking at the charts

Indications for surgery

Contact surgical colleagues at the outset, before an operation is necessary rather than when it is inevitable. Delay increases mortality. When the following criteria are met, surgery may be appropriate. Any decision not to operate should only be taken after discussion with the consulting surgeon.

- Age > 60 years and
 - > 4 units transfused in 24 h, or
 - · one rebleed in hospital, or
 - · continued bleeding, or
 - · spurting vessel at endoscopy
- Age < 60 years and
 - > 8 units transfused in 24 h, or
 - · one rebleed in hospital, or
 - · continued bleeding, or
 - spurting vessel at endoscopy

The differential diagnosis of upper gastrointestinal bleeding is shown in Table 1.1 (p. 10). Individual topics are discussed below.

Oesophageal varices

Cirrhosis is the commonest cause of portal hypertension (p. 152) and oesophageal varices in the UK and North America. Whilst oesophageal varices can be found at endoscopy in almost 50% of patients with cirrhosis, less than one-third of these will bleed from their varices. Mortality during an acute bleed depends on the severity of liver disease on admission. Mortality according to Child's grade A is 10%, grade B is 25% and grade C is 50% (Table 5.5, p. 144), but 60–80% of all patients who bleed from varices will be dead within 4 years.

Assessment

- Bleeding from oesophageal varices is a complex clinical emergency for which control of bleeding is only one aspect
- Attention to infection, control of ascites (p. 147), encephalopathy (p. 144), alcohol withdrawal (Fig. 5.8, p. 177) and nutrition (p. 371) are vital for a successful outcome

Acute bleeding

• Resuscitate and monitor (p. 11). Colloids (synthetic, albumin or blood) are indicated and saline can be used in the acute situation. Avoid 5% dextrose if hyponatraemic

- 30% with known varices have another source of haemorrhage
- During active bleeding, correct disordered coagulation to international normalized ratio (INR) < 1.5 or prothrombin time < 22 s with fresh frozen plasma (FFP). However, FFP is contraindicated in the absence of bleeding, because this increases intravascular volume and variceal pressure, which may precipitate haemorrhage. Platelet transfusion may be necessary for thrombocytopenia (platelet count < 60 × 10⁹/L). Discuss with the haematologists
- Arrange urgent endoscopy for banding or sclerotherapy by an experienced endoscopist
- Control of the airway, with endotracheal intubation if necessary, is extremely important when carrying out emergency endoscopy in a patient with an active variceal bleeding
- Give oral lactulose, starting at 90 mL/day or phosphate enemas if nil by mouth to prevent/treat hepatic encephalopathy (p. 145), intravenous vitamins (p. 393) and benzodiazepine, as necessary, for alcohol withdrawal (p. 177).

If endoscopic therapy is not available, or massive bleeding continues:

- Insert a Sengstaken tube until sclerotherapy/banding can be performed or repeated after 12 h
- Start splanchnic vasoconstrictors such as Glypressin (2 mg bolus, then 2 mg every 4 h, for up to 96 h). Glypressin rarely can be associated with ischaemic complications
- For recurrent bleeding after two attempts at endoscopic therapy the alternatives are transjugular intrahepatic portosystemic shunt (TIPSS) or oesophageal transection (see below)
- Where a stent is inserted between the hepatic and portal vein under radiological control, TIPSS can only be performed by experienced interventional radiologists, but is probably the procedure of choice where bleeding is not controlled by other means (Section 5.4, p. 153). Control of bleeding is excellent (approaching 100%), but 1 month mortality is still high (30–40%) owing to liver failure. Late complications (blocked stent, encephalopathy) are common
- Oesophageal transection has a mortality of 50% and should only be considered for
 patients without other organ failure, who were admitted in Child's group A or B
 (Table 5.5, p. 144). Splenectomy and proximal gastric devascularisation are needed to
 prevent subsequent bleeding from gastric varices
- Bleeding gastric varices are one cause of failed endoscopic haemostasis. TIPSS should be considered (Section 5.4, p 153), but preliminary reports suggest that endoscopic injection of bovine thrombin (2–10 mL of 1000 U/mL) or histoacryl (mixed with 1:1 lipiodal, 1–2 mL injected) controls active bleeding, although varices are not eradicated. Histoacryl has the potential to glue up the endoscope and should only be used by experienced endoscopists

Balloon tamponade

- Indicated for uncontrolled variceal bleeding, or recurrent haemorrhage despite sclerotherapy or banding
- To be inserted by experienced operators only
- Sedation, or a general anaesthetic to insert an endotracheal tube and secure the airway, may be necessary
- Insert a cooled, lubricated Sengstaken or Minnesota tube beyond 45 cm. The tube can usually be stiffened by inserting a well-lubricated pair of paediatric endoscopic biopsy forceps down the central lumen

- Inflate the gastric balloon with 300 mL tap water containing 50 mL of any intravenous X-ray contrast medium or 300 mL of air. Ensure gastric balloon channel is double-clamped
- Tie a 250 mL bag of saline to the tube to provide traction at the gastrooesophageal junction, but be very careful to protect the mouth to prevent pressure necrosis
- Aspirate gastric and oesophageal ports (if present) hourly, and connect to a bag for continuous drainage
- It is rarely necessary to inflate the oesophageal balloon. If necessary because of persistent bleeding, inflate the oesophageal balloon to 30 mmHg with air, measured by manometer. Deflate for 5 min every hour.
- X-ray to check position (Fig. 1.2)
- Active bleeding is arrested in 90%. Continued bleeding usually means that the tube is misplaced or that there are gastric varices
- Deflate the oesophageal balloon after 6 h and the gastric balloon after 12–24 h to allow further endoscopic sclerotherapy or banding
- Complications include tracheal intubation, oesophageal rupture from inflating the gastric balloon in the oesophagus, or mucosal necrosis from leaving the balloon inflated for too long

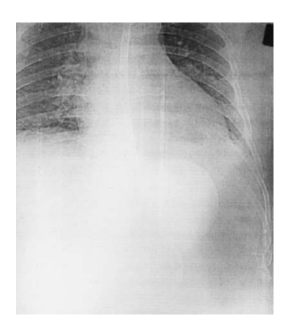
Prevention of rebleeding from varices

- The highest risk of rebleeding is in the first 6 weeks
- Most gastroenterologists repeat endoscopic therapy at 1–2-week intervals until varices are obliterated
- Varices recur after obliteration in 40%, usually within 1 year
- Propranolol 40 mg three times daily decreases the risk of bleeding, can be used following initial endoscopic therapy and in this situation is as effective as repeated endoscopic therapy.
- Portal hypertensive gastropathy may cause bleeding after variceal obliteration. The gastric mucosa (usually fundal) has a characteristic 'snakeskin' appearance at endoscopy. Propranolol is the treatment of choice (p. 153)
- Surgical or percutaneous TIPSS are indicated for recurrent bleeding, especially in non-cirrhotic portal hypertension (p. 153)

Mallory–Weiss tears

A mucosal tear at the oesophagogastric junction due to forceful vomiting results in haematemesis. The typical features are:

- · Initial vomitus does not contain blood
- Vomiting has often been provoked by alcohol
- 90% settle with conservative treatment
- Acid-suppressing drugs are unnecessary
- Continued bleeding can be controlled by endoscopic injection or thermocoagulation. Surgery is very rarely needed
- Bleeding after forceful vomiting may also be caused by prolapse of the gastric mucosa resulting in a focal area of haemorrhagic gastropathy ('hernia gastropathy') opposite the gastro-oesophageal junction



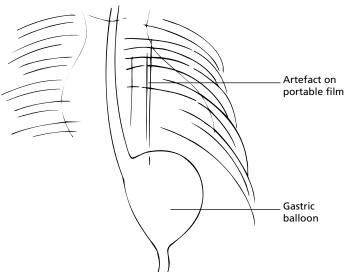


Fig. 1.2 X-ray showing Minnesota tube in correct position. Gastric balloon has been inflated with 300 mL water mixed with 50 mL contrast medium.

Acute gastric erosions and haemorrhagic gastropathy

Erosions are diagnosed endoscopically, but may be obscured by oozing from haemorrhagic gastropathy (gastritis is a misleading term that should be reserved for a histological diagnosis). Major haemorrhage from superficial gastric injury is unusual and the cause is often readily apparent.

1.5 Acute bleeding: upper gastrointestinal tract

Causes

- Non-steroidal anti-inflammatory drugs (NSAIDs)
- · Alcohol
- Stress (trauma, major surgery or patients in intensive care)

Specific treatment

- A PPI (any) is usually given for 1–4 weeks, depending on the cause
- Persistent bleeding is treated with intravenous tranexamic acid 1 g three times daily, in addition to a PPI (e.g. omeprazole or pantoprazole), which can be given intravenously
- Total gastrectomy is the last resort for continued bleeding after all medical treatment
 has been vigorously applied for 24–48 h, and should only be performed by an experienced surgeon
- It should be noted that despite the fact that antacids, ranitidine or sucralfate may prevent stress erosions in intensive care patients, there is almost no evidence that they reduce clinically significant bleeding or mortality

Gastric ulcer (Section 3.5, p. 92)

- Consider provoking causes (such as NSAIDs)
- Appropriate endoscopic haemostasis (p. 12) reduces rebleeding and mortality
- Give a PPI (lansoprazole 30 mg, omeprazole 20 mg, pantoprazole 40 mg, rabeprazole 20 mg daily) for 4 weeks once bleeding has stopped, together with *H. pylori* eradication therapy if endoscopic biopsies confirm infection
- For patients with NSAID-associated ulcers who cannot stop NSAIDs (p. 109), concomitant PPI therapy (lansoprazole or omeprazole) has replaced misoprostol in healing ulcers and preventing recurrence
- Arrange a repeat endoscopy after 8–12 weeks, to biopsy and take brushings for cytology from the ulcer site
- If surgery is needed for continued bleeding, a Billroth 1 gastrectomy is usually performed. Undersewing with a vagotomy and pyloroplasty is a simpler operation, but the ulcer cannot be examined histologically to exclude cancer. Wedge resection removes the ulcer and has the lowest morbidity in high-risk elderly patients, but long-term acid suppression is then necessary because it does not prevent recurrent ulceration

Duodenal ulcer (Section 3.9, p. 102)

- Combine ulcer healing with eradication of H. pylori
- The optimum treatment is triple, or quadruple eradication therapy (Table 3.2, p. 89)
- Repeat endoscopy is unnecessary, except in special circumstances (e.g. patients needing warfarin)
- Always confirm that eradication of H. pylori has been successful after an ulcer has bled, preferably by an isotope breath test (p. 86)
- Successful eradication of *H. pylori* significantly reduces the risk of rebleeding and the risk of bleeding from another ulcer to an extent similar to acid suppression. It is also cheaper in the long term and provides a cure
- Risk of repeat haemorrhage without eradication or maintenance therapy is 20% over 5–10 years, but is higher if associated with NSAIDs
- NSAID-associated ulcers heal with PPIs even if NSAIDs have to be continued, but are not directly associated with *H. pylori* (p. 110)

1.5 Acute bleeding: upper gastrointestinal tract

Maintenance acid suppression (lansoprazole 15 mg, omeprazole 10 mg daily) is only
indicated for patients at high risk of dying from the complications of recurrent ulceration (p. 110) when eradication therapy has been unsuccessful, or when NSAIDs have
to be continued

'No source of bleeding found'

This is common (up to 20% acute bleeding) and can produce difficult management problems. Possible causes are:

- · Lesion missed on endoscopy
- Mucosal lesion healed before patient endoscoped:
 - · erosions
 - · Mallory-Weiss tear
 - Dieulafoy's lesion (bleeding vessel with no surrounding ulceration, usually high on the greater curve)
- Bleeding from third part of the duodenum, or beyond:
 - jejunum (ulcerative jejunitis)
 - · Meckel's diverticulum
 - colon
- · Other:
 - · nose bleed
 - rare causes of bleeding (Table 1.1, p. 10)

Management

The management of gastrointestinal bleeding from obscure and occult sources is discussed in more detail on p. 8 and Section 9.7 (p. 329; Fig. 9.9, p. 330).

- Reassess the patient—no further action is necessary for low-risk patients (p. 9)
- Repeat endoscopy in patients with a predicted mortality > 10% (p. 9)
- Investigate rare causes of bleeding (recheck coagulation, discuss small bowel radiology, endoscopic retrograde cholangiopancreatography (ERCP), ⁹⁹Tc sulphur colloid red cell scan (p. 20) or ⁹⁹Tc pertechnate scan with radiologists)
- Selective angiography during active bleeding (which must be at a rate of 1 U/4 h) is indicated after two negative endoscopies, preferably in a specialist unit
- Small bowel enteroscopy, including video capsule endoscopy (Section 9.7, p. 329) may be available in specialist units, but referral is necessary, and other procedures (mesenteric angiography, small bowel enema) will normally be repeated in the specialist unit
- Laparotomy, careful examination of the whole bowel with a bright light (e.g. sigmoi-doscopy light source) and peroperative endoscopy is the ultimate procedure for recurrent episodes of active bleeding from obscure origin, but may still not identify the source and it is usually wise to refer to a specialist unit if this is contemplated

Aortoenteric fistula

Consider this rare diagnosis in *every* patient with an aortic graft and gastrointestinal bleeding. Exsanguination at the first bleed is uncommon. Small 'herald' bleeds occur for up to 2 weeks. Urgent abdominal CT scan, the diagnostic procedure of choice, may show haematoma around the graft. Endoscopy, if performed, should be to the fourth part of the duodenum, but surgery should not be delayed if hypotension has occurred. Aggressive surgery is needed as soon as the diagnosis is made, preferably in a specialist unit.