Bowel Obstruction

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Introduction
Acute intestinal obstruction accounts for 10% of emergency surgical admissions\(^1\). This occurs when there is an interruption to the forward flow of intestinal contents. Intestinal obstruction is associated with complications such as ischaemia, perforation and aspiration pneumonia. The decision making process can be quite challenging; surgeons have to make critical decisions with regard to non-operative management vs surgery. Delays in diagnosis, radiological investigation and surgical intervention can significantly affect patient outcome. Adhesions from previous surgery are currently the leading cause of small bowel obstruction in industrialized countries (~70%), followed by malignancy, inflammatory bowel disease, and hernias\(^2\). Malignancy and volvulus are the commonest causes of large bowel obstruction.

According to Hwang et al\(^3\), small bowel obstruction alone accounts for more than 300,000 operations annually in the United States\(^2\). Most of the patients managed with bowel obstruction require attention to nutrition, electrolyte balance, symptom control and intestinal decompression. At the moment there is no national guideline or framework for the management of acute intestinal obstruction. There are emerging guidelines for the management of small and large bowel obstruction, however it is felt that there is considerable variation of the management of bowel obstruction, with variation in outcomes\(^4, 5, 6\).

Bowel obstruction may be caused by intrinsic or extrinsic mechanical obstruction. Patients with small bowel / high intestinal obstruction are likely to experience symptoms of vomiting and central colicky abdominal pain. Patients with large bowel/low intestinal obstruction are likely to experience the symptoms of abdominal distension and constipation\(^7\).

Non-operative management of all forms of bowel obstruction includes good initial resuscitation with oxygen and IV fluids, NG tube aspiration, adequate analgesia and close monitoring of the patient for signs of ischaemia or perforation, when early surgical intervention would be needed. This includes regular clinical assessment for peritonitis and observation monitoring for signs of sepsis.

Specific non-operative therapeutic intervention for adhesive small bowel obstruction includes a water soluble contrast study which has a diagnostic and therapeutic value and may overcome the obstruction by drawing fluid into the bowel (osmotic effect) resulting in a shorter hospital stay\(^8\).

For large bowel obstruction due to tumours colonic stenting may be used as a definitive treatment or as a bridge to surgery\(^9, 10, 11, 12\). This is not currently widely adopted across the UK as a treatment option and anecdotally there is significant variation in its use.

Surgery for small bowel obstruction may involve laparotomy and division of adhesions, small bowel resection, stoma formation, hernia repair and bypass operations. Increasing use of laparoscopic surgery for small bowel obstruction has been seen but consensus has not been reached as to the indications for this treatment\(^13\).

Large bowel obstruction may be treated surgically by bowel resection, stoma or bypass.

When surgery is required, mortality can exceed 10%, far higher that seen in elective colorectal surgery. The majority of patients requiring surgery can be categorized as “high-risk” and require consultant delivered care and admission to critical care after surgery.
Prompt recognition of patient deterioration, sepsis, and perforation is needed. Surgery may be required within a matter of hours for surgical source control of sepsis, or to prevent impending perforation.  

**Guidelines and standards**

NASBO: Report of the National Audit of Small Bowel Obstruction, 2017


Association of Surgeons of Great Britain and Northern Ireland, Royal College of Surgeons Commissioning guide: Emergency general surgery (acute abdominal pain), 2014


ACPGI Emergency General Surgery Sub-Committee. Recommendations for the management of large bowel obstruction, June 2017


Aims and objectives

Overall aim:
To identify remedial factors in process of care of patients with both large and small intestinal obstruction.

Objectives

Clinical
To identify and assess the care delivered to patients presenting with acute bowel obstruction at all points in the pathway including:

- Emergency admission factors (including recognition prior to hospital admission)
- Initial assessment and diagnosis (including risk assessment and delays in diagnosis)
- Admission to the ward (including the route of admission, admitting specialty and delays in admission)
- Treatment plan (including continuity of care and communication)
- Imaging (including time to imaging, the reporting of imaging and communication of results)
- Decision making (including multidisciplinary input and clinician seniority)
- Non-surgical therapy
- Nutrition
- Surgery (including delays, decision making and continuity of care)
- Post operative care (including location, nutrition and complications)
- Discharge and follow up arrangements
- End of life care

Organisational
To examine organisational aspects of care including:

- Access to services (including CT, multisite working and emergency surgery)
- Staffing
- The use of local and national guidelines and protocols
- Networks of care
- Staff training regarding acute bowel obstruction management
- Communication
- Audit/governance

Methods

Population/Inclusions
Data will be collected on all patients aged 16 and over admitted to hospital with an obstructed bowel over a four week period.

Patients will be identified using the following ICD10 codes (ICD10 code for acute bowel obstruction to be recorded in the first three positions in order for the patient to be included in the study)

- K56.1 Intussusception
- K56.2 Volvulus
- K56.3 Gallstone ileus
- K56.4 Other impaction of intestine
- K56.5 Intestinal adhesions [bands] with obstruction
- K56.6 Other and unspecified intestinal obstruction
- K40.0 Bilateral inguinal hernia, with obstruction, without gangrene
K40.1 Bilateral inguinal hernia, with gangrene
K40.3 Unilateral or unspecified inguinal hernia, with obstruction, without gangrene
K40.4 Unilateral or unspecified inguinal hernia, with gangrene
K41.0 Bilateral femoral hernia, with obstruction, without gangrene
K41.1 Bilateral femoral hernia, with gangrene
K41.3 Unilateral or unspecified femoral hernia, with obstruction, without gangrene
K41.4 Unilateral or unspecified femoral hernia, with gangrene
K42.0 Umbilical hernia with obstruction, without gangrene
K42.1 Umbilical hernia with gangrene
K43.0 Ventral hernia with obstruction, without gangrene
K43.1 Ventral hernia with gangrene
K44.0 Diaphragmatic hernia with obstruction, without gangrene
K44.1 Diaphragmatic hernia with gangrene
K45.0 Other specified abdominal hernia with obstruction, without gangrene
K45.1 Other specified abdominal hernia with gangrene
K46.0 Unspecified abdominal hernia with obstruction, without gangrene
K46.1 Unspecified abdominal hernia with gangrene

Exclusions
The following will be excluded where recorded in isolation (i.e. not with an included ICD10 code):
- K55 Vascular disorders of the intestine
- K56.0 Paralytic ileus
- K56.7 Ileus, unspecified
- K91.3 Postoperative intestinal obstruction

Participating sites
All hospital providers that admit/treat patients with bowel obstruction including: acute hospitals, specialist hospitals, community hospitals, independent hospitals, treatment centres etc will be asked to participate in the study.

Sample size
Based on HES data, over a one year period (2016/17) there were 55,996 patients admitted to hospital with one of the included ICD10 codes; this equates to approximately 1077 admissions per week.

Based on data collected as part of the pilot study, there were an average of 333 cases per hospital per year, which equates to just over 6 cases per hospital per week. If data were collected from 300 hospitals over a 4 week period (there are approx. 340 ‘acute’ sites listed on our DB) this would give us a sample of approximately 7200 cases.

Within the pilot data, approximately 6% of patients died prior to discharge, and 25% of patients underwent surgery (so this would be approximately 432 deaths within the pool, and 1800 patients who underwent surgery)

- Sample size for clinician questionnaire – 2000 cases
• Sample for case note review – To be confirmed once the assessment form has been designed.

If we need to sample cases for inclusion, this will be based on including:
• All deaths
• A sample of patients who underwent surgery
• A sample of patients (no surgery/discharged alive) + AKI (N17) and/or ventilation (E85.1 & E85.2)

**Sample period**
Data will be collected over a four week period from 00:00 Monday 16th April– 23:59 Sunday 13th May 2018.

**Case identification**
Within each Trust/Health Board NCEPOD has a Local Reporter (usually employed in clinical audit) who is responsible for providing the details of cases for inclusion to NCEPOD. At the start of the study the Local Reporter will be contacted and sent details of the study criteria. Patients with bowel obstruction will be identified retrospectively through ICD10 coding via completion of a spreadsheet with selected data from central hospital records. This will include patient details (NHS number, hospital number, age), admission/discharge dates, patient destination (including death)/source, ICD10 codes (primary and all, relating to bowel obstruction and AKI), details of the admitting consultant, operation details (responsible consultant, dates and OPCS codes), OPCS codes for any ventilation provided, and details of any previous admissions for bowel obstruction in the previous 12 months.

**Method of data collection**
**Clinician questionnaire**
A clinical questionnaire will be sent to the consultant who was responsible for the patient at the time of hospital admission. Within this there will be instruction to pass the questionnaire on to most appropriate clinician (i.e. to the operating clinician if the patient underwent the surgery). The questionnaire will be sent to the NCEPOD local reporter for dissemination. Reminder letters will be sent at six weeks and ten weeks where the data are outstanding.

**Case notes for peer review**
• Clinical notes from the time of admission to discharge
• Critical care notes
• Care pathways
• Nursing notes
• Fluid balance charts
• Weight charts
• Malnutrition Universal Screening Tool (MUST) score
• Food charts
• Oral care charts
• Radiology and xray reports
• Endoscopy reports
• Biochemistry and haematology reports
• Theatre notes
• Anaesthetic charts
• Consent forms
• Drug charts
• Discharge summary
• Any separate dietetics notes

Upon receipt at NCEPOD the case notes will be made anonymous for patient identifiable information.

**Reviewer Assessment form:**
A multidisciplinary group of reviewers (detailed below) will assess the case notes and clinician questionnaires at review meetings held at the NCEPOD office and give their opinion on quality of care via completion of the reviewer assessment form.

**Organisational questionnaire**
An organisational questionnaire will be disseminated to all participating sites and collect data on organisational aspects of care of patients with bowel obstruction.

**Pilot Study**
A pilot study will be performed to test the method of data collection (including the feasibility of accessing critical care and primary care data) and the data collection materials and ensure that they are robust.

**Analysis and Review of Data**

**Case Reviewers**
A multidisciplinary group of reviewers will be recruited to review the data collected and provide opinion on the care received by this group of patients, from admission to discharge. The advisor group would be made up of surgeons (including general, gastrointestinal and colorectal surgeons), radiologists, nurses, physicians (including emergency medicine and acute physicians), intensivists, anaesthetists, and dieticians.

**Data Entry**
All clinician questionnaire data will be electronically collected and combined with data from the assessment form completed by the case reviewers. Quantitative data analysis will be undertaken using Excel and qualitative analysis will be undertaken by reviewing the themes arising from the Advisor meetings.

**Confidentiality and data protection**
Once the data have been extracted by the NCEPOD researchers, the questionnaires and casenotes will be anonymised to remove patient identifiers prior to review by the case reviewers. All electronic data are held in password protected files and all paper documents in locked filing cabinets. As soon as possible after receipt of data NCEPOD will encrypt electronic identifiers and anonymise paper documents. Section 251 approval has been obtained to perform this study without the use of patient consent.

**Dissemination**
On completion of the study a report will be published and widely disseminated.

**Timescale**

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**References**


