# A randomised controlled trial to assess whether the amount of mesentery removed, or the type of bowel join used during surgery for Crohn's disease, can affect the chances of getting further disease

Submission date	Recruitment status Recruiting	[X] Prospectively registered		
16/02/2022		[X] Protocol		
Registration date	Overall study status	[X] Statistical analysis plan		
24/02/2022	Ongoing  Condition category	Results		
Last Edited		☐ Individual participant data		
09/12/2025	Digestive System	[X] Record updated in last year		

### Plain English summary of protocol

Background and study aims

Crohn's is a disease that makes the bowel red, swollen and painful. It is a lifelong disease. Some people get few flare-ups. Other people may have really bad and long-lasting symptoms. There is no cure for Crohn's, but drugs can treat the symptoms. If the drugs do not work surgery may be needed. Whilst surgery can also stop symptoms for a long time, the disease can return. Further drugs and even further surgery may be needed. Many surgeons feel that the way the bowel and the tissue containing the blood and other vessels supplying the bowel (the mesentery) is removed and the way the healthy bowel ends are re-joined can affect the success of surgery. Two changes to the way surgeons operate have been proposed. One involves taking out more of the mesentery and the other involves doing another type of bowel join. This is known as the Kono-S join. They are safe but researchers need to test that they are better than the usual method in stopping further disease. In this study the researchers will ask people if they would let them carry out one or both of these methods. They will then see if they have reduced the chances of further disease 1 year later. The type of surgery a person has will be decided by chance.

### Who can participate?

Patients who are undergoing surgery (ileocaecal resection) for Crohn's disease where the bowel is re-joined (anastomosis).

### What does the study involve?

If a patient is eligible and happy to be part of the study, they will be asked to sign a consent form. They will then complete three questionnaires and be asked some questions about their medical history. On the day of the surgery the surgeon will confirm the patient is eligible and the patient will be allocated to one of the study groups by chance (randomised). The surgeon will then perform the type of surgery as decided by the randomisation. Six weeks after surgery

patients will be asked to complete the questionnaires again during their visit to the hospital (this visit is part of standard care for all patients having this kind of surgery, even if they are not in the trial). After 6 to 12 months patients normally have a colonoscopy as part of the follow-up after the surgery. At this time the participants will be asked to complete the three questionnaires again. The researchers will also look at the participants' medical notes for up to 3 years after the surgery to see if Crohn's disease has come back. At some sites participants will have additional blood and tissue samples taken during their surgery and colonoscopy. These will be analysed to help the researchers understand what might be causing Crohn's disease.

What are the possible benefits and risks of participating?

Taking part in this study will mean no additional appointments at the hospital compared to the number of appointments patients receive normally. The risks associated with the different surgery types are not thought to be any different to usual surgery for Crohn's disease, and the surgeon will discuss this further with patients as part of standard surgery pre-assessment. Several studies have shown that all the surgical procedures are safe. Complications that are associated with Crohn's surgery will have been discussed with the patient and include: leak of the join in the bowel, bleeding, bowel blockage (ileus), wound infection, urinary infection, heart events, clots in the leg or lungs, chest infection, hernia, and damage to other parts of the abdomen. Additionally, very rarely patients may have a reaction to the anaesthetic.

Where is the study run from? The University of Sheffield (UK)

When is the study starting and how long is it expected to run for? September 2021 to July 2027

Who is funding the study? National Institute for Health Research (NIHR) Efficacy and Mechanism Evaluation (EME) Programme (UK)

Who is the main contact? Steven Brown, steven.brown13@nhs.net

## Contact information

## Type(s)

Scientific

#### Contact name

Dr Jamie Hall

#### **ORCID ID**

https://orcid.org/0000-0003-4042-5591

#### Contact details

Room 2.15, The Innovation Centre c/o Regent Court 30 Regent Street Sheffield United Kingdom S1 4DA

## Additional identifiers

### Clinical Trials Information System (CTIS)

Nil known

### Integrated Research Application System (IRAS)

301301

### ClinicalTrials.gov (NCT)

Nil known

### Central Portfolio Management System (CPMS)

52048

## Study information

### Scientific Title

MEsenteric Excision and Kono-S Anastomosis Trial (MEErKAT)

### Acronym

**MEErKAT** 

### **Study objectives**

- 1. How and why should surgeons consider the mesentery and anastomotic technique in Crohn's disease?
- 2. Does Kono-S anastomosis reduce the recurrence of Crohn's disease over standard anastomosis?
- 3. Does radical mesenteric resection reduce the recurrence of Crohn's disease over standard mesenteric resection?
- 4. Does a combination of these techniques reduce the recurrence of Crohn's disease over standard mesenteric resection?

### Ethics approval required

Old ethics approval format

## Ethics approval(s)

Approved 05/04/2022, North East - Tyne & Wear South Research Ethics Committee (+44 (0) 2071048306, +44 (0)22071048285, +44 (0)22071048265; tyneandwearsouth.rec@hra.nhs.uk), REC Ref: 22/NE/0041

## Study design

Randomized; Interventional; Design type: Treatment, Surgery

## Primary study design

Interventional

## Study type(s)

### **Treatment**

### Health condition(s) or problem(s) studied

Crohn's disease

#### **Interventions**

The trial follows the standard pathway for ileocaecal Crohn's. Participants are recruited when surgery is deemed appropriate. Potential participants will be discussed at MDT meetings and identified at the time of pre-operative assessment before the day of surgery. Surgeons, research nurses and trainees will also check waiting lists for those listed for ileocaecal resection for primary or recurrent Crohn's disease. If a patient is interested in taking part, the local research team will review information about their Crohn's disease from their medical records.

A member of the patient's care team will identify and consent eligible participants. Potential participants will receive an approved Participant Information Sheet (PIS) and be given the opportunity to ask questions from both the surgical and research team. Potential participants will be approached either at their clinic visit prior to surgery or pre-operative assessment. At a pre-operative assessment or on the day of surgery, the surgeon, research nurse or trainee will offer entry into the trial and consent will be obtained. No study-related procedures will occur before the approved consent form is signed, other than the initial case note review for eligibility.

A screening log will be maintained for each hospital site, to document all potential participants screened, whether they were recruited, and any reasons for non-recruitment where this information is available. Screening logs will be requested and reviewed by CTRU on a regular basis.

#### Baseline Visit

If a patient is happy to proceed, and their surgeon feels the trial is appropriate for them, they will be asked to sign a consent form to join the trial. Following consent, they will then complete separate questionnaires about how they feel (physically and mentally) and about their Crohn's disease. These are called the IBD-Control, CDAI and EQ-5D questionnaires. These are to help researchers understand a patient's health at the start of the trial and will ask them general questions about their health and specific questions about their Crohn's disease. The researchers will also take details of the patient's medical history, personal demographics (e.g. gender, ethnicity, etc.) and any medications they are taking.

### Day of Surgery

On the patient's day of surgery (which could be the same as the baseline visit), their surgeon will confirm that they still qualify for the trial and they will be randomised to one of the four combinations of surgical techniques (the University of Sheffield provides the randomisation system). The treatment options are as follows:

There are two groups of mesenteric excision and two groups of anastomosis:

Mesenteric Excision Groups

Radical Mesenteric Excision

More mesentery is removed along with the diseased bowel, whilst preserving the main blood supply to the remaining bowel

Close Mesenteric Excision

Minimal mesentery is removed, staying close to the bowel wall.

### **Anastomosis Groups**

Patients randomised to Kono-S

For this anastomosis the resected bowel is stapled perpendicular to the mesentery and the stapled ends sutured together to form the supporting column. The bowel is joined together by hand using stitches and is a different way to join the bowel.

### Patients randomised to Standard of Care

Standard care is essentially surgeons' preference of anastomosis. Anastomosis may utilise staples or sutures to join the small bowel and colon after the diseased bowel is removed.

Essentially patients will be randomised onto one of 4 combinations in the ratio 1:1:1:1:

- 1. Kono-S + radical mesenteric resection
- 2. Kono-S + close mesenteric resection
- 3. Standard anastomosis + radical mesenteric resection
- 4. Standard anastomosis + close mesenteric resection or surgeons' choice

For all cases the mode of access (laparoscopic or open), closure technique and post-operative care are according to usual practice for that participating centre. Whichever surgical technique is selected, it will not have a detrimental effect for patients in the event further surgical procedures are required if the disease does reoccur at the site of resection. The treatment group will not be revealed to the patients (they are blinded), and so are the endoscopists (who perform colonoscopy) at the follow-up visits. It is usual practice for surgeons to take part of the bowel out during the operation and send this for microscopic analysis. The researchers will take small samples of the bowel that has been removed for additional analysis, to review as part of the trial also. They need to see why these new surgical methods may work and clues may be found by looking at those who get further disease after surgery and seeing what part of the join the disease has come back to. They will also leave a small tattoo in the inside of the patient's remaining bowel so that we can identify it when they come back for a telescope assessment of the bowel. This telescopic assessment (colonoscopy) is always carried out 6-12 months after operation, whether patients are in the trial or not. These tattoos are used all the time when doing telescopic camera tests and know they are safe. The tattoo will only be seen if someone looks into the bowel. Patient's care after surgery will be the same as people have after this surgery and who are not in the trial.

It is also useful for our research to take pictures of the bowel that has been removed after surgery has been completed. The researchers will ask patients to consent to having a photo taken of their removed tissue after surgery, directly after the procedure, and will be of the removed tissue pinned onto a specimen board only. They will also take a photograph of the bowel at their follow up colonoscopy to help look at disease recurrence. The anonymised photos will be uploaded to the secure server database at the University of Sheffield that complies with the Data Protection Act (2018) and uses industry-standard techniques to provide security. The photos will be used for the central assessment of each surgery, to ensure consistency in surgical technique across the trial and disease assessment.

#### Week 6

After the patient has recovered from their surgery and has been discharged from hospital, they will come back to clinic at around 6 weeks after the procedure. This is in line with standard of care after this type of surgery. At this visit, the researchers will check how they are and they will also be asked to complete the IBD-Control, CDAI and EQ-5D questionnaires again.

### Follow Up

As mentioned earlier, it is usual after this operation to have a colonoscopy about 6 to 12 months

after surgery. This looks at the bowel join (including the tattoo inked at surgery). It is also usual to have some small biopsies taken at this time. The research team will also ask patients to complete the three questionnaires - IBD-Control, CDAI and EQ-5D – and ask what medications they are taking and whether they have had any side effects since their operation. Finally, they will also look at patients' medical notes up to three years after their surgery to see if any recurrence has occurred. The trial does not involve any other visits or treatments above normal routine care.

### Mechanistic Sub-Study

The researchers are running a small optional sub-study for a group of around 140 patients to assess what might be causing recurrent Crohn's disease. It is referred to as a 'mechanistic substudy' as it is looking at the mechanisms (e.g. types of tissue, immune cells, etc.) of why Crohn's disease may reoccur in more detail, along with data from the main trial. If a patient agrees to take part in this part of the study, an additional set of blood and tissue samples will be taken at their surgery and colonoscopy follow up:

Immediately before surgery:

Blood samples (5-10ml, around 2 teaspoons)

### At operation:

Extra tissue samples 5-10mm in size will be taken from the part of the bowel and mesentery that has been removed.

### At colonoscopy:

2-4 tiny biopsies will be taken from the lining of the small bowel

Blood samples (5-10 ml, around two teaspoons):

No extra visits are required, as the samples will be taken at the existing trial visits. Consenting to this part of the trial is entirely optional for patients and will only be run at specific centres.

### Intervention Type

Procedure/Surgery

### Primary outcome(s)

Time to endoscopic recurrence of disease using the Modified Rutgeerts score (≥i2) from the date of randomisation after 6-12 months to a maximum of 3 years follow-up

## Key secondary outcome(s))

- 1. Incidence of severe endoscopic recurrence measured using the Modified Rutgeerts score ≥i3 after 6-12 months to a maximum of 3 years follow-up
- 2. Clinician and patient-reported symptomatic recurrence at 6-12 months and at the end of the trial. The definition of symptomatic recurrence will be:
- 2.1. Self-reported recurrence in combination with endoscopically confirmed recurrence; OR
- 2.2. Surgical re-intervention in combination with histological confirmation of recurrence; OR
- 2.3. Change of medical strategy for reasons other than safety/tolerability; OR
- 2.4. IBD Control >13; OR, Crohn's Disease Activity Index (CDAI) >220
- 3. Quality of life measured using EQ-5D-5L at 6-12 months after surgery
- 4. Time to endoscopic recurrence for all groups measured using Modified Rutgeerts score (≥i2) for at least 6 months and up to 3 years after surgery
- 5. Surgical recurrence measured using clinician and patient report at a minimum of 6 months and a maximum of 3 years

6. Radiological and surgical anastomotic leak (following surgery) as defined by the latest consensus; other complications for each intervention (surgical complications may include, but are not limited to haemorrhage; ileus/bowel obstruction; wound infection; urinary tract infection; cardiac events; pulmonary embolism [PE]/deep vein thrombosis [DVT]; and respiratory insufficiency/pneumonia. Late postoperative complications may include: trocar-site and incisional hernia; ureteral stenosis [retroperitoneal fibrosis]) recorded at surgery, following surgery and within at least 6 months and a maximum of 3 years following surgery 7. Mesenteric disease severity measured using the mesenteric disease activity index at surgery

### Completion date

30/07/2027

## **Eligibility**

### Key inclusion criteria

People aged over 18 years undergoing ileocaecal resection for primary/recurrent Crohn's disease where an anastomosis is carried out

### Participant type(s)

**Patient** 

### Healthy volunteers allowed

No

### Age group

Mixed

### Lower age limit

18 years

### Upper age limit

999 years

### Sex

All

### Total final enrolment

0

### Key exclusion criteria

Participant exclusion criteria as of 10/11/2023:

- 1. Patients with markedly extensive inflammation affecting the vascular root of the mesentery seen on imaging or at operation
- 2. Patients undergoing stoma formation proximal to the anastomosis
- 3. Patients who have a contraindication to subsequent colonoscopy
- 4. Patients unable to give full informed consent
- 5. Patients who are pregnant (as ascertained by standard pregnancy tests undertaken at preoperative visits as per standard clinical care)
- 6. Patients who, in the opinion of the principal investigator, do not meet the criteria for relevant surgery

Previous participant exclusion criteria:

- 1. Patients with markedly extensive inflammation affecting the vascular root of the mesentery seen on imaging or at operation
- 2. Patients undergoing stoma formation
- 3. Patients who have a contraindication to subsequent colonoscopy
- 4. Patients unable to give full informed consent
- 5. Patients who are pregnant (as ascertained by standard pregnancy tests undertaken at preoperative visits as per standard clinical care)
- 6. Patients who, in the opinion of the principal investigator, do not meet the criteria for relevant surgery

# Date of first enrolment 02/05/2022

Date of final enrolment 31/01/2026

## Locations

### Countries of recruitment

United Kingdom

England

Scotland

## Study participating centre Wythenshawe Hospital

Southmoor Road Wythenshawe Manchester England M23 9LT

## Study participating centre Northern General Hospital

Northern General Hospital NHS Trust C Floor, Huntsmnan Building Herries Road Sheffield England S5 7AU

## Study participating centre

### Norfolk & Norwich University Hospital

Colney Lane Colney Norwich England NR4 7UY

## Study participating centre Guys Hospital

Great Maze Pond London England SE1 9RT

## Study participating centre Churchill Hospital

Old Road Headington Oxford England OX3 7LE

# Study participating centre University Hospital (coventry)

Clifford Bridge Road Coventry England CV2 2DX

## Study participating centre Queen Elizabeth Hospital

Mindelsohn Way Edgbaston Birmingham England B15 2GW

# Study participating centre Solihull Hospital

Lode Lane Solihull England B91 2JL

## Study participating centre Royal Cornwall Hospital (treliske)

Treliske Truro England TR1 3LJ

## Study participating centre Russells Hall Hospital

Pensnett Road Dudley England DY1 2HQ

## Study participating centre Manchester Royal Infirmary

Cobbett House Oxford Road Manchester England M13 9WL

# Study participating centre Bristol Royal Infirmary

Marlborough Street Bristol England BS2 8HW

## Study participating centre Central Middlesex Hospital

Acton Lane London England NW10 7NS

## Study participating centre Kettering General Hospital

Rothwell Road Kettering England NN16 8UZ

## Study participating centre Leicester Royal Infirmary

Infirmary Square Leicester England LE1 5WW

## Study participating centre Huddersfield Royal Infirmary

Acre Street Huddersfield England HD3 3EA

## Study participating centre Chelsea & Westminster Hospital

369 Fulham Road London England SW10 9NH

## Study participating centre Bolton Royal Hospital

Minerva Road Farnworth Bolton England BL4 0JR

## Study participating centre Queen's Hospital Burton

Belvedere Road

Burton on Trent England DE13 0RB

## Study participating centre Queen Elizabeth University Hospital

1345 Govan Road Glasgow Scotland G51 4TF

# Study participating centre Derriford Hospital

Derriford Road Derriford Plymouth England PL6 8DH

## Study participating centre Royal Devon and Exeter Hospital

Royal Devon & Exeter Hospital Barrack Road Exeter England EX2 5DW

## Study participating centre Sandwell District General Hospital

Lyndon West Bromwich England B71 4HJ

## Study participating centre Northern General Hospital

Northern General Hospital NHS Trust C Floor, Huntsmnan Building Herries Road Sheffield England S5 7AU

## Study participating centre Southampton General Hospital

Tremona Road Southampton England SO16 6YD

## Study participating centre Royal Albert Edward Infirmary

Wigan Lane Wigan England WN1 2NN

## Study participating centre Northumbria Healthcare NHS Foundation Trust

North Tyneside General Hospital Rake Lane North Shields England NE29 8NH

## Study participating centre Royal Alexandra Hospital

Corsebar Road Paisley Scotland PA2 9PN

## Sponsor information

### Organisation

Sheffield Teaching Hospitals NHS Foundation Trust

**ROR** 

## Funder(s)

## Funder type

Government

### Funder Name

NIHR Evaluation, Trials and Studies Co-ordinating Centre (NETSCC); Grant Codes: NIHR131988

## **Results and Publications**

### Individual participant data (IPD) sharing plan

The results will be published on a freely accessible database within 1 year of completion of the trial.

## IPD sharing plan summary

Stored in publicly available repository

## **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
HRA research summary			28/06/2023		No
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes
Protocol file	version 2.0	28/06/2023	19/01/2024	No	No
Statistical Analysis Plan	version 1.0	24/07/2023	19/01/2024	No	No
Study website	Study website	11/11/2025	11/11/2025	No	Yes