Testing for bile duct stones before gallbladder surgery

Submission date	Recruitment status	[X] Prospectively registered		
24/09/2018	No longer recruiting	[X] Protocol		
Registration date	Overall study status	Statistical analysis plan		
02/10/2018	Ongoing	Results		
Last Edited	Condition category Surgery	Individual participant data		
13/12/2024		[X] Record updated in last year		

Plain English summary of protocol

Background and study aims

Surgery to remove the gallbladder is required if it contains gallstones that cause problems. About 70,000 operations are performed annually in England. Sometimes, gallstones cause other problems if they pass from the gallbladder into the nearby bile duct (e.g. jaundice/inflammation of the pancreas). If this happens, it is necessary to remove the bile duct stones before or during the gallbladder operation. Because of this, patients requiring gallbladder surgery are assessed for any risk of bile duct stones. If the risk is high, further tests are done to identify if bile duct stones are present. If the risk is moderate or low (although it can be difficult to distinguish between the two), then it is uncertain whether further tests to look for bile duct stones are necessary. As a result, some surgeons choose to perform tests and others don't. A UK-wide research study found that a third of patients undergoing gallbladder surgery were tested, usually before surgery using an MRI scanner. This test, called a Magnetic Resonance Cholangiopancreatography (MRCP), involves a 1-hour visit to hospital and costs the NHS about £365. The MRCP identifies bile duct stones but may delay gallbladder surgery which can lead to increased problems with gallstones whilst waiting. There are other uncertainties about the need for testing using MRCP. Even if the MRCP shows bile duct stones, the stones can pass into the bowel spontaneously; and removing the stones can cause complications. Not having the MRCP avoids these risks, but can lead to bile duct stones being left behind after surgery, which may also cause complications. Research to establish if going straight to gallbladder surgery without testing the bile duct beforehand is needed.

The Sunflower Study will find out whether testing for bile duct stones with MRCP before gallbladder surgery is worthwhile or not in patients with a low or moderate risk of having stones.

Who can participate?

Adults with symptomatic gallstone disease who are scheduled and fit to receive gallbladder surgery, with a low to moderate risk of common bile duct stones

What does the study involve?

Participants in the Sunflower study will be divided into two groups. One group will go straight to surgery (i.e. no additional testing) and the other will receive an MRCP before surgery. The groups will be selected by a process called randomisation to ensure that groups have similar patients in terms of factors such as general health, age and gender. The 'straight to surgery'

group will have twice as many people in as the 'tested' group to reduce the number of extra MRCPs performed. Both groups will be followed for 18 months and information about the need for treatment of bile duct stones, complications of surgery and costs collected.

What are the possible benefits and risks of participating?

For participants who receive MRCP, the possible benefit of participating is that this procedure may detect and, if needed, treat problematic bile duct stones. However for participants who do not receive MRCP, the possible benefit is that this allows their gallbladder surgery to go ahead immediately, without the possible disadvantages of testing for bile duct stones. For participants who receive MRCP, the possible risks of participating is that the procedure may

be unnecessary, as bile duct stones often pass safely and spontaneously into the bowel. Additionally, in 10-20% of cases, MRCP may not detec bile duct stones. Additionally, some patients may experience claustrophobia during the scan. For some patients, the scan and endoscope procedure (if required) may lead to a longer wait time for gallbladder surgery, which can lead to problems with gallstones whilst waiting (however, the surgery may not always be delayed by this).

For participants who do not receive MRCP, the possible risk is that any bile duct stones present will not be detected, which could lead to problems after gallbladder surgery (i.e. jaundice, infection, pancreatitis) and therefore require further treatment or readmission to hospital. If bile duct stones are suspected later, a scan and potentially endoscope procedure will be likely needed to remove them.

Where is the study run from? Leeds Teaching Hospitals NHS Trust and at least 50 hospitals throughout the UK.

When is the study starting and how long is it expected to run for? January 2018 to November 2025

Who is funding the study? National Institute for Health Research (NIHR) Health Technology Assessment Programme (UK)

Who is the main contact? Stephen Palmer sunflower-study@bristol.ac.uk

Contact information

Type(s)
Public

Contact name

Mr Stephen Palmer

Contact details

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Additional identifiers

Clinical Trials Information System (CTIS)

Nil known

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

16/142/04

Study information

Scientific Title

The Sunflower Study: A randomised controlled trial to establish the clinical and cost effectiveness of expectant management versus pre-operative imaging with MRCP in patients with symptomatic gallstones undergoing laparoscopic cholecystectomy at low or moderate risk of common bile duct stones

Acronym

The Sunflower Study

Study objectives

We will test the hypothesis that expectant management is non-inferior to magnetic resonance pancreaticogram (MRCP) with respect to hospitalisation for treatment for a complication of gallstones up to 18 months after randomisation.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved 10/12/2018, Yorkshire & The Humber – South Yorkshire Ethics Committee (NHSBT Newcastle Blood Donor Centre, Holland Drive, Newcastle upon Tyne, NE2 4NQ; Tel: +44 (0)207 1048091; Email: nrescommittee.yorkandhumber-southyorks@nhs.net), ref: 18/YH/0358

Study design

Interventional multi-centre pragmatic randomized controlled trial with an internal pilot phase (Phase

I) and a quintet recruitment intervention (QRI)

Primary study design

Interventional

Study type(s)

Diagnostic

Health condition(s) or problem(s) studied

Symptomatic gallbladder disease, requiring gallbladder surgery

Interventions

Participants will be randomly allocated in a 1:2 ratio to receive either a reoperative magnetic resonance pancreaticogram (magentic resonance cholangiopancreatography - MRCP) or no MRCP (expectant management) using an online system.

In the MRCP arm, participants will undergo an MRCP prior to their gallbladder surgery. An MRCP takes on average 20-30 minutes. In the expectant management arm, participants will proceed directly to surgery without undergoing an MRCP. The follow up will be the same in both groups – a 20% sample will be asked to complete questionnaires at 3, 6, 12 and 18 months post randomisation.

Intervention Type

Other

Primary outcome(s)

Current primary outcome measure:

Any of the following:

- 1. Any hospital admission within 18 months of randomisation for treatment of complications of gallstones whether in the CBD or gallbladder
- 2. Complications during the admission for LC for the treatment for gallstones or any readmission for complications of the LC leading to a hospital stay of >2 days. Complications will include, but not be limited to:
- 2.1. Return to theatre post LC for any cause
- 2.2. Percutaneous radiological drainage
- 2.3. ERCP for non-diagnostic reasons (e.g. for a bile leak). It does not include a diagnostic ERCP performed following an MRCP where CBD stones were identified.
- 3. Complications during any ERCP for the treatment for gallstones. Complications will include:
- 3.1. Blood transfusion post ERCP
- 3.2. Percutaneous radiological drainage
- 3.3. Treatment of a perforation occurring during ERCP
- 3.4. Acute pancreatitis
- 3.5. Other complications leading to a hospital stay of >2 days

This will be acquired from routine data sources, such as HES data from NHS Digital in England. The HES data will be reviewed and specific OPCS-4 and ICD-10 codes will be searched for. These codes identify specific diagnoses, procedures, etc which will form the primary outcome measure. This will be measured in the 18 months following randomisation.

Previous primary outcome measure:

Any of the following:

- 1. Any hospital admission within 18 months of randomisation for treatment of complications of gallstones whether in the CBD or gallbladder
- 2. Complications during the admission for LC for the treatment for gallstones or readmission for complications of the LC. Complications comprise:
- 2.1. Return to theatre post LC for any cause
- 2.2. Percutaneous radiological drainage
- 2.3. ERC
- 3. Complications during the index admission for ERCP for the treatment for gallstones (i.e. an ERCP performed following an MRCP where CBD stones were identified). Complications comprise:
- 3.1. Blood transfusion post ERCP

- 3.2. Percutaneous radiological drainage
- 3.3. Further ERCP
- 3.4. Treatment of a perforation occurring during ERCP
- 3.5. Acute pancreatitis
- 3.6. Respiratory complications leading to extended hospital stay of > 2 days
- 3.7. Cardiac complications leading to extended hospital stay of > 2 days
- 3.8. Infective complications leading to extended hospital stay of > 2 days
- 3.9. Other complications leading to extended hospital stay of > 2 days (detail to be recorded). This will be acquired from routine data sources, such as HES data from NHS Digital in England. The HES data will be reviewed and specific OPCS-4 and ICD-10 codes will be searched for. These codes identify specific diagnoses, procedures, etc which will form the primary outcome measure. This will be measured in the 18 months following randomisation.

Key secondary outcome(s))

- 1. Health-related quality of life, assessed using the EQ-5D-5L questionnaire at time of randomisation, admission for LC and 3, 6, 12 and 18 months after randomisation (participants will not be asked to complete the admission for LC questionnaire if they have completed their baseline questionnaire within the previous two days)
- 2. Items in the LC core outcome set, which is due to be published at the end of 2018
- 3. NHS resource use to 18 months post randomisation, taken from routine data sources such as HES data from NHS Digital in England

Completion date

30/11/2025

Eligibility

Key inclusion criteria

Current inclusion criteria as of 15/01/2020:

- 1. Aged 18 years or older
- 2. Symptomatic gallbladder disease, confirmed by trans-abdominal ultrasound scan (USS) or computed tomography (CT) scan, including, for example:
- 2.1. Biliary colic
- 2.2. Cholecystitis
- 2.3. Mild and severe gallstone pancreatitis
- 2.4. Gallbladder polyps
- 2.5. Gallbladder dyskinesia. etc
- 3. Scheduled and fit for laparoscopic cholecystectomy (LC) as an elective or urgent procedure
- 4. Low or moderate risk of common bile duct (CBD) stones, including all of the following:
- 4.1. CBD diameter ≤8 mm on USS
- 4.2. Bilirubin ≤ 50umol/l
- 4.3. Alanine transferase less than three times the upper limit of normal ($\leq 3 \times ULN$) and/or alkaline phosphatase $\leq 3 \times ULN$

If a patient does not meet the definition of low or moderate risk of CBD stones solely because both alanine transferase and alkaline phosphatase are $> 3 \times ULN$, if repeat blood tests are carried out and at least one of the second or subsequent test results is within range (i.e. $\le 3 \times ULN$) the patient may be recruited at that time.

If a patient does not meet the definition of low or moderate risk of CBD stones solely because bilirubin > 50umol/l, if repeat blood tests are carried out and at least one of the second or

subsequent test results is within range the patient may be recruited at that time. If CBD cannot be seen on USS or CT scan, the patient may be recruited as long as all the other inclusion criteria are met and there is no intrahepatic duct dilatation reported.

Previous inclusion criteria:

- 1. Aged 18 years or older
- 2. Symptomatic gallstone disease, confirmed by trans-abdominal ultrasound scan (USS) or computed tomography (CT) scan, including:
- 2.1. Biliary colic
- 2.2. Cholecystitis
- 2.3. Mild and severe pancreatitis
- 2.4. Gallbladder polyps
- 2.5. Gallbladder dyskinesia
- 3. Scheduled and fit for laparascopic cholecystectomy (LC) as an elective or urgent procedure
- 4. Low or moderate risk of common bile duct (CBD) stones, including all of the following:
- 4.1. CBD diameter ≤8 mm on USS
- 4.2. Bilirubin ≤ 50umol/l
- 4.3. Alanine transferase less than twice the upper limit of normal ($\leq 2 \times ULN$) and/or alkaline phosphatase $\leq 2 \times ULN$.

If a patient does not meet the definition of low or moderate risk of CBD stones solely because both alanine transferase and alkaline phosphatase are $> 2 \times ULN$, if repeat blood tests are carried out and at least one of the second or subsequent test results is within range (i.e. $\le 2 \times ULN$) the patient may be recruited at that time.

If a patient does not meet the definition of low or moderate risk of CBD stones solely because bilirubin > 50umol/l, if repeat blood tests are carried out and at least one of the second or subsequent test results is within range the patient may be recruited at that time.

If CBD cannot be seen on USS or CT scan, the patient may be recruited as long as all the other inclusion criteria are met and there is no intrahepatic duct dilatation reported.

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

Αll

Total final enrolment

7471

Key exclusion criteria

Current exclusion criteria as of 15/01/2020:

- 1. Unable to undergo MRCP
- 2. Evidence of empyema or perforated gallbladder requiring urgent intervention

- 3. Previous gastric bypass
- 4. Previous MRCP or endoscopic ultrasound (EUS) within last 3 months
- 5. Any previous ERCP
- 6. Haemolytic disease
- 7. Pregnancy
- 8. Unwilling to participate in follow up
- 9. Unable to provide written informed consent
- 10. Prisoner

Previous exclusion criteria:

- 1. Unable to undergo MRCP
- 2. Evidence of empyema or perforated gallbladder requiring urgent intervention
- 3. Previous duodenal bypass
- 4. Previous MRCP within last 3 months
- 5. Haemolytic disease
- 6. Pregnancy
- 7. Unwilling to participate in follow up
- 8. Unable to provide written informed consent
- 9. Prisoner

Date of first enrolment

01/12/2018

Date of final enrolment

30/08/2024

Locations

Countries of recruitment

United Kingdom

England

Northern Ireland

Scotland

Wales

Study participating centre Leeds Teaching Hospitals NHS Trust United Kingdom LS1 3EX

Study participating centre

Calderdale and Huddersfield NHS Foundation Trust

Huddersfield Royal Infirmary

Acre Street Lindley Huddersfield United Kingdom HD3 3EA

Study participating centre Nottingham University Hospitals NHS Trust

Queens Medical Centre Campus Derby Road Nottingham United Kingdom NG7 2UH

Study participating centre Worcestershire Acute Hospitals NHS Trust

Worcestershire Royal Hospital Worcester United Kingdom WR5 1DD

Study participating centre North Bristol NHS Trust

Southmead Hospital Bristol United Kingdom BS10 5NB

Study participating centre University Hospitals Bristol NHS Foundation Trust

Marlborough Street Bristol United Kingdom BS2 8HW

Study participating centre Surrey and Sussex Healthcare NHS Trust

East Surrey Hospital Canada Avenue Redhill United Kingdom RH1 5RH

Study participating centre Countess of Chester Hospital NHS Foundation Trust

Countess of Chester Hospital Liverpool Road Chester United Kingdom CH2 1UL

Study participating centre Buckinghamshire Healthcare NHS Trust

Stoke Mandeville Hospital Mandeville Road Aylesbury United Kingdom HP21 8AL

Study participating centre Northumbria Healthcare NHS Foundation Trust

North Tyneside General Hospital Rake Lane North Shields United Kingdom NE29 8NH

Study participating centre University Hospitals Plymouth NHS Trust

Derriford Hospital Plymouth United Kingdom PL6 8DH

Study participating centre Northern Devon Healthcare NHS Trust

North Devon District Hospital Raleigh Park Barnstaple United Kingdom EX31 4JB

Study participating centre Guy's and St Thomas' NHS Foundation Trust

St Thomas' Hospital Westminster Bridge Road London United Kingdom SE1 7EH

Study participating centre Sherwood Forest Hospitals NHS Foundation Trust

Kings Mill Hospital Mansfield Road Sutton In Ashfield United Kingdom NG17 4JL

Study participating centre Great Western Hospitals NHS Foundation Trust

Great Western Hospital Marlborough Road Swindon United Kingdom SN3 6BB

Study participating centre Sheffield Teaching Hospital NHS Foundation Trust

Northern General Hospital Sheffield United Kingdom S5 7AU

Study participating centre Basildon and Thurrock University Hospitals NHS Foundation Trust Basildon University Hospital Nethermayne

Basildon United Kingdom SS16 5NL

Study participating centre University Hospital of Derby and Burton NHS Foundation Trust

Queen's Hospital Belvedere Road Burton-on-Trent United Kingdom DE13 0RB

Study participating centre Gloucestershire Hospitals NHS Foundation Trust

Gloucester Royal Hospital Great Western Road Gloucester United Kingdom GL1 3NN

Study participating centre Bradford Teaching Hospitals NHS Foundation Trust

Bradford Royal Infirmary Duckworth Lane Bradford United Kingdom BD9 6RJ

Study participating centre

County Durham and Darlington NHS Foundation Trust

University Hospital of North Durham North Road Durham United Kingdom DH1 5TW

Study participating centre

The Royal Bournemouth and Christchurch Hospitals NHS Foundation Trust

Royal Bournemouth Hospital Castle Lane East Bournemouth United Kingdom BH7 7DW

Study participating centre Royal Free NHS Trust

Pond St London United Kingdom NW3 2QG

Study participating centre Royal Devon and Exeter Foundation Trust Hospital

Barrack Road Exeter United Kingdom EX2 5DW

Study participating centre Hereford County Hospital

Hereford United Kingdom HR1 2ER

Study participating centre Royal Bolton Hospital

Minerva Road Bolton United Kingdom BL4 0JR

Study participating centre Royal Blackburn Hospital

Haslingden Road Blackburn United Kingdom BB2 3HH

Study participating centre

Tameside General Hospital

Fountain Street Ashton Under Lyne United Kingdom OL6 9RW

Study participating centre The Whittington Hospital

Magdala Avenue London United Kingdom N19 5NF

Study participating centre Royal Derby Hospital

Uttoxeter Rd Derby United Kingdom DE22 3NE

Study participating centre Musgrove Park Hospital

Taunton United Kingdom TA1 5DA

Study participating centre Worthing Hospital

Lyndhurst Road Worthing United Kingdom BN11 2DH

Study participating centre Royal Albert and Edward Infirmary

Wigan Lane Wigan United Kingdom WN1 2NN

Study participating centre Weston General Hospital

Grange Road Uphill Weston-Super-Mare United Kingdom BS22 4TQ

Study participating centre Royal Lancaster Infirmary

Ashton Road Lancaster United Kingdom LA1 4RP

Study participating centre St Mary's Hospital

Parkhurst Road Newport United Kingdom PO30 5TG

Study participating centre University Hospitals Coventry and Warwickshire

Clifford Bridge Road Coventry United Kingdom CV2 2DX

Study participating centre Forth Valley Royal Hospital

Stirling Road Larbert United Kingdom FK5 4WR

Study participating centre

University Hospital of North Tees

Stockton-on-Tees United Kingdom TS19 8PE

Study participating centre Southend University Hospital NHS Foundation Trust

Westcliff on Sea United Kingdom SS0 0RY

Study participating centre Sunderland Royal Hospital

Kayll Road Sunderland United Kingdom SR4 7TP

Study participating centre Royal United Hospitals Bath NHS Foundation Trust

Combe Park Bath United Kingdom BA1 3NG

Study participating centre Aneurin Bevan University Health Board

Royal Gwent Hospital Cardiff Road Newport United Kingdom NP20 2UB

Study participating centre St George's Hospital NHS Foundation Trust

Ground Floor Jenner Wing Cranmer Terrace London United Kingdom SW17 ORE

Study participating centre Milton Keynes University Hospital

Standing Way Eaglestone Milton Keynes United Kingdom MK6 5LD

Study participating centre University Hospitals Birmingham NHS Foundation Trust Birmingham United Kingdom B15 2TH

Study participating centre University Hospital Southampton NHS Foundation Trust Southampton General Hospital Southampton United Kingdom SO16 6YD

Study participating centre Doncaster Royal Infirmary Armthorpe Road Doncaster United Kingdom DN2 5LT

Study participating centre Freeman Hospital Newcastle Upon Tyne United Kingdom NE7 7DN

Study participating centre
Northampton General Hospital
Cliftonville
Northampton

United Kingdom NN5 1BD

Study participating centre Belfast Health and Social Care Trust

Trust Headquarters, A Floor Belfast City Hospital 51 Lisburn Road Belfast United Kingdom BT9 7AB

Study participating centre York and Scarborough Teaching Hospitals NHS Foundation Trust

Wiggington Road York United Kingdom YO31 8HE

Study participating centre Queen Elizabeth Hospital Gateshead

Queen Elizabeth Avenue Gateshead United Kingdom NE9 6SX

Study participating centre St Mary's Hospital

London United Kingdom W2 1NY

Study participating centre Princess of Wales Hospital

Coity Road Bridgend Bridgend County Borough United Kingdom CF31 1RQ

Study participating centre Hull University Teaching Hospitals NHS Trust

Castle Hill Hospital Castle Road Cottingham United Kingdom HU16 5JQ

Study participating centre Ipswich Hospital

Heath Road Ipswich United Kingdom IP4 5PD

Study participating centre Shrewsbury & Telford Hospital NHS Trust

Mytton Oak Road Shrewsbury United Kingdom SY3 8XQ

Sponsor information

Organisation

Leeds Teaching Hospitals NHS Trust

ROR

https://ror.org/00v4dac24

Funder(s)

Funder type

Not defined

Funder Name

Health Technology Assessment Programme

Alternative Name(s)

NIHR Health Technology Assessment Programme, Health Technology Assessment (HTA), HTA

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location

United Kingdom

Results and Publications

Individual participant data (IPD) sharing plan

Data requests would be sent to the study manager via sunflower-study@bristol.ac.uk. The full de-identified dataset would be available and will be held indefinitely. Access would be dependent on provision of an ethically approval study protocol and could cover numerous analysis types. The mechanism of data sharing would be determined at the time. Consent from participants will be obtained for data sharing. Data will be de-identified. Ethical approval for a new project would need to be in place.

IPD sharing plan summary

Available on request

Study outputs

Output type Details	Date created	Date l added	Peer reviewed?	Patient- facing?
Protocol article	29/06 /2021	13/08 /2021	Yes	No
HRA research summary		28/06 /2023	No	No
Participant information sheet version 7.0	11/07 /2022	29/07 /2022	No	Yes
Participant information sheet sheet	11/11 /2025	11/11 /2025	No	Yes
Protocol file version v4.0	18/08 /2020	23/09 /2020	No	No
Protocol file version 5.0	08/09 /2021	27/10 /2021	No	No
Protocol file version 7.0	07/09 /2023	17/10 /2023	No	No
13/12/2024: The sunflower-study contact stated that based on funder recommendations protocol v8.0 had been withdrawn and the withdrawal has been confirmed by the HRA/REC, which fully withdrew Protocol file the amendment to introduce version 8.0. Protocol v7.0, dated 07/09 /2023 is in use. version 8.0	10/10 /2024	20/11 /2024	No	No
Study Study website website	11/11 /2025	11/11 /2025	No	Yes