

# Pilonidal sinus treatment: studying the options

<b>Submission date</b> 03/12/2018	<b>Recruitment status</b> No longer recruiting	<input checked="" type="checkbox"/> Prospectively registered <input type="checkbox"/> Protocol
<b>Registration date</b> 26/03/2019	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input checked="" type="checkbox"/> Results
<b>Last Edited</b> 26/09/2024	<b>Condition category</b> Infections and Infestations	<input type="checkbox"/> Individual participant data

## Plain English summary of protocol

### Background and study aims

Pilonidal disease is an infection in the crease of a person's buttocks. The aim of this study is to determine the types of pilonidal disease, to describe the various treatments, to engage with patients and to determine which outcomes they value and which treatments they prefer/do not prefer, and to provide recommendations for further research.

### Who can participate?

Patients aged over 16 years with symptomatic pilonidal sinus

### What does the study involve?

Before the start of the study there will be a survey circulated within the colorectal consultant surgeon network. This survey will seek to determine the methods of surgery currently employed in the field and the ways in which the surgeon learned this technique. The survey will also explore the factors affecting choice of treatment method in relation to disease presentation. Finally, it will seek to investigate the estimated mean recurrence rate for each surgeon. The study will record details of participants' pit/track anatomy, surgical treatment, medical events, and health-related quality of life until six months after their operation. It will then describe the combination of treatments currently in use and quantify clinical and patient-reported outcomes associated with each. It will also identify patient-specific disease features that might predict poor outcomes in each treatment group. It will then derive an estimate of the risks associated with common treatment options, ranking the best treatment strategies among patients for whom more than one treatment is considered appropriate. The study will then use case-studies to provide an overview of patient views and experiences of the treatment that they have chosen and received. This will also include the views of patients on which treatments they would rather avoid and which outcomes they most value in regards to their treatment options.

### What are the possible benefits and risks of participating?

The study is predicted to reach a consensus on which subtypes of pilonidal disease may benefit from which treatment options. Additionally, the research is predicted to reach a surgeon and patient-based consensus on research priorities.

### Where is the study run from?

University of Sheffield and 28 sites (UK)

When is the study starting and how long is it expected to run for?  
September 2018 to March 2023

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

Who is the main contact?  
1. Emily Strong  
EmilyStrong@barnsley.gov.uk  
2. Prof. Steven Brown  
3. Prof Daniel Hind  
d.hind@sheffield.ac.uk

## Contact information

### Type(s)

Public

### Contact name

Ms Emily Strong

### ORCID ID

<https://orcid.org/0000-0002-2381-4088>

### Contact details

ScHARR  
University of Sheffield  
Regent Court  
30 Regent Street  
Sheffield  
United Kingdom  
S1 4DA  
+44 (0)114 2222974  
EmilyStrong@barnsley.gov.uk

### Type(s)

Scientific

### Contact name

Prof Steven Brown

### ORCID ID

<https://orcid.org/0000-0002-0980-2793>

### Contact details

Northern General Hospital  
Herries Rd  
Sheffield  
United Kingdom  
S5 7AU

**Type(s)**

Scientific

**Contact name**

Prof Daniel Hind

**ORCID ID**

<https://orcid.org/0000-0002-6409-4793>

**Contact details**

ScHARR

University of Sheffield

Regent Court

30 Regent Street

Sheffield

United Kingdom

S1 4DA

-

d.hind@sheffield.ac.uk

**Additional identifiers****Clinical Trials Information System (CTIS)**

Nil known

**Protocol serial number**

CPMS 40221

**Study information****Scientific Title**

The PITSTOP study: pilonidal sinus treatment: studying the options

**Acronym**

PITSTOP

**Study objectives**

This is a prospective cohort study to determine the subtypes of pilonidal disease for which the various treatment options are indicated, to describe the various interventions that are used to treat the disease, to engage with patients and determine which outcomes they value and which interventions they prefer/do not prefer and to provide recommendations for further research.

**Ethics approval required**

Old ethics approval format

**Ethics approval(s)**

East of England - Cambridge South Research Ethics Committee

The Old Chapel

Royal Standard Place  
Nottingham  
NG1 6FS

26/11/2018, ref: 18/EE/0370

## **Study design**

Observational; Design type: Cohort study

## **Primary study design**

Observational

## **Study type(s)**

Treatment

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

Pilonidal sinus disease

## **Interventions**

Design

1. Survey of consultant surgeons
2. Observational cohort
3. Nested mixed method case study
4. Discrete choice experiment
5. Modified nominal group consensus exercise

### **1. 1. Survey of consultant surgeons**

Prior to the commencement of the cohort study there will be a survey circulated within the colorectal consultant surgeon network. The survey will be distributed via email throughout the trainee surgeon network and will be completed in paper and pencil format by consultant colorectal surgeons. This survey will seek to determine the methods of surgery currently employed in the field and the ways in which the surgeon learned this technique. The survey will also explore the factors affecting choice of treatment method in relation to disease presentation. Finally, it will seek to investigate the estimated mean recurrence rate for each surgeon. Currently, there is a lack of evidence outlining the general practice of treatment for pilonidal sinus in the UK. This survey will provide evidence on the techniques currently in use in the UK Health Care system.

### **2. Observational cohort study**

Patients considered suitable for surgery will be identified from GP or secondary care referrals, sent a patient information sheet about the study, detailing the study and all treatment options, and diverted preferably to a specific recruiting clinic. At the clinic, a member of the research team will explain the study to the patient and give them the opportunity to ask any questions. The Principal Investigator or delegated research team member will confirm eligibility and ensure written informed consent is obtained before any patient data is collected. Participants will be advised that they are able to withdraw from the study at any point without any impact on their routine NHS care. As is standard practice, the surgeon will discuss the condition, possible interventions and their advantages and disadvantages. Each participant will be invited to answer a questionnaire on shared decision making (CollaboRATE).

Baseline data, taken face to face immediately after consent, will include patient demographics, social and occupation factors, hair and skin type and previous pilonidal surgery history. Pilonidal disease characteristics, recorded by the surgeon immediately post-op will include pit numbers, track numbers, length, unilateral/bilateral distribution, position, presence of pus and previous surgical scarring. Description of the intervention will include all excision and closure techniques used with a description of major steps taken and potential variants thereof. Use of anaesthetic (general, regional, local), period-operative antibiotics, planned post-operative wound care will also be recorded.

A pain numeric rating scale score will be recorded by the patient on day 1 via text reminder. A trained research nurse will collect structured questionnaire data and a pain numeric rating scale by telephone at seven days and six months, but face-to-face at the routine clinic visit at 4-6 weeks post-operatively.

Hospital notes and patient self-report will be used to assess recurrence and complications at six months. A patient regret score will also be collected at 6 months.

Following the six-month assessment all patients will be contacted with a one-off follow-up at the end of the study unless the participant withdraws consent to further follow up or dies before the study completes. Patients will be contacted via email and telephone with a questionnaire incorporating persistence of symptoms. Hospital and GP records will be used to ascertain A&E attendance, repeat/additional procedures and unresolved complications due to PD, and interactions with primary care (GP consultations for PD, practice or district nurse visits for dressings).

For all of the cohort patients, data will be collected to establish which patients have further treatment for recurrent symptoms or complications following their initial procedure. This will be achieved at the six-week clinic visit following the intervention and by interrogating hospital records, asking the patients' consultants, writing to patients' GPs and questioning the patient via telephone interview at 6 months and at the end of the study. At this stage each participant will also be invited to complete a questionnaire on decision regret.

### 3. Nested mixed method case study

A proportion of patients (at least 20) will be randomly selected using a theoretically based sample and asked to take part in the nested mixed methods substudy. For the participants' comfort and convenience, we will collect data by telephone or Skype. To assess which outcomes are most valued by patients, and whether there are particular interventions they would rather avoid, the baseline interview guide will adapt key questions from the CODE framework. Interviews will last 15-20 minutes at baseline and 10 minutes at six months.

### 4. Discrete choice experiment

The data from these interviews will be used to construct a discrete choice experiment. All consented patients will be sent a link to their email addresses via Qualtrics, containing a participant information sheet and simple instructions on how to complete the choice questionnaire. The questionnaire will contain hypothetical choice scenarios (typically range from 8-12 choice sets to avoid cognitive burden) and will ask patients to make choices between two combinations of outcomes with varying levels. The questionnaire should take no more than 15 minutes to complete.

### 5. Modified nominal group consensus exercise

At the end of the cohort study a consensus regarding the sub-groups of patients for whom the various interventions may be suited will be generated, along with a consensus working with

clinicians and patients, together with a consensus meeting defining appropriate comparators and valued outcomes for any future randomised controlled trial. To do this, a modified nominal group technique consensus exercise will be undertaken. This will take place over half a day, adjacent to the annual ACPGBI. It will be opened to around 40 colorectal surgeons and around 15 patients from across the UK.

## **Intervention Type**

Other

## **Primary outcome(s)**

As this is a cohort study examining current practice there are no primary outcome measures per se. Identifying appropriate outcome measures for future studies is one of the objectives of the project. The trialists aim to collect the following data:

1. Pain measured using a numeric rating scale at day 1 and day 7 post-operatively. It will also be collected at the routine follow up appointment and at the six month follow up
2. Quality of life measured using the EQ-5D-5L quality of life scale at baseline, at day 7 post operative, at the routine follow up appointment and at the six month follow up
3. Interactions with primary and secondary care asked at day 7 post-operatively, at the routine follow up appointment and at the six month follow up
4. Wound healing and impact measured using the Cardiff Wound Impact Scale at the routine follow up appointment and at the six month follow up
5. Return to normal activities asked using a single question at at baseline, at day 7 post operative, at the routine follow up appointment and at the six month follow up
6. Complications that occur following the intervention identified on the 'Procedure details' CRF and any further complications identified at the six-week clinic visit and at the six-month follow-up
7. Recurrence measured through a single question at an end of study follow up call or through hospital notes

## **Key secondary outcome(s)**

There are no secondary outcome measures

## **Completion date**

31/03/2023

# **Eligibility**

## **Key inclusion criteria**

Current inclusion criteria as of 04/09/2020:

1. Consenting patients with pilonidal sinus disease that requires surgical intervention
2. Aged over 16 years

Previous inclusion criteria:

1. Consenting patients with pilonidal sinus disease that requires surgical intervention
2. Aged over 18 years

## **Participant type(s)**

Patient

## **Healthy volunteers allowed**

No

**Age group**

Adult

**Lower age limit**

16 years

**Sex**

All

**Total final enrolment**

677

**Key exclusion criteria**

1. Asymptomatic
2. Pregnant
3. Unable to give consent
4. Acute abscess - which is defined as a tender swelling containing pus that presents acutely - the decision on whether or not to operate is at the discretion of the surgeon assessing the patient

**Date of first enrolment**

01/05/2019

**Date of final enrolment**

31/03/2022

**Locations****Countries of recruitment**

United Kingdom

England

Scotland

Wales

**Study participating centre**

**Addenbrooke's Hospital**

Hills Rd

Cambridge

United Kingdom

CB2 0QQ

**Study participating centre**

**University Hospital of Wales**

Heath Park Way

Cardiff  
United Kingdom  
CF14 4XW

**Study participating centre**  
**Countess of Chester Hospital**  
Liverpool Rd  
Chester  
United Kingdom  
CH2 1UL

**Study participating centre**  
**Royal Derby Hospital**  
Uttoxeter Rd  
Derby  
United Kingdom  
DE22 3NE

**Study participating centre**  
**Glasgow Royal Infirmary**  
84 Castle St  
Glasgow  
United Kingdom  
G4 0SF

**Study participating centre**  
**Leicester General Hospital**  
Gwendolen Rd  
Leicester  
United Kingdom  
LE5 4PW

**Study participating centre**  
**Wirral University Teaching Hospital NHS Foundation Trust (WUTH)**  
Arrowe Park Hospital  
Arrowe Park Rd  
Birkenhead  
United Kingdom  
CH49 5PE

**Study participating centre**  
**Manchester Royal Infirmary**  
Oxford Rd  
Manchester  
United Kingdom  
M13 9WL

**Study participating centre**  
**Norfolk and Norwich University Hospitals NHS Foundation Trust**  
Colney Ln  
Norwich  
United Kingdom  
NR4 7UY

**Study participating centre**  
**John Radcliffe Hospital**  
Headley Way  
Headington  
Oxford  
United Kingdom  
OX3 9DU

**Study participating centre**  
**Queen Alexandra Hospital**  
-  
Portsmouth  
United Kingdom  
PO6 3LY

**Study participating centre**  
**St. Mark's Hospital**  
Watford Rd  
Harrow  
United Kingdom  
HA1 3UJ

**Study participating centre**  
**Northern General Hospital**  
Glossop Rd  
Sheffield

United Kingdom  
S10 2JF

**Study participating centre**

**Western General Hospital**

Crewe Rd S  
Edinburgh  
United Kingdom  
EH4 2XU

**Study participating centre**

**Tameside and Glossop Integrated Care NHS Foundation Trust**

Fountain St  
Ashton-under-Lyne  
United Kingdom  
OL6 9RW

**Study participating centre**

**Newcastle Upon Tyne Hospitals NHS Foundation Trust**

Freeman Road  
High Heaton  
Newcastle upon Tyne  
United Kingdom  
NE7 7DN

**Study participating centre**

**York Teaching Hospital NHS Foundation Trust**

Wigginton Road  
York  
United Kingdom  
YO31 8HE

**Study participating centre**

**St Helens and Knowsley Teaching Hospitals NHS Trust**

Whiston Hospital  
Warrington Road  
Prescot  
United Kingdom  
L35 5DR

**Study participating centre**

**Morrison Hospital**

Heol Maes Eglwys  
Morrison  
Cwmrhydyceirw  
Swansea  
United Kingdom  
SA6 6NL

**Study participating centre**

**Salford Royal NHS Foundation Trust**

Stott Ln  
Salford  
United Kingdom  
M6 8HD

**Study participating centre**

**Yeovil District Hospital NHS Foundation Trust**

Higher Kingston  
Yeovil  
United Kingdom  
BA21 4AT

**Study participating centre**

**Musgrove Park Hospital**

Parkfield Dr  
Taunton  
United Kingdom  
TA1 5DA

**Study participating centre**

**Queen Elizabeth The Queen Mother Hospital**

St Peter's Rd  
Margate  
United Kingdom  
CT9 4AN

**Study participating centre**

**Royal Devon & Exeter NHS Foundation Trust**

Exeter

United Kingdom  
EX2 5DW

**Study participating centre**  
**East Suffolk and North Essex NHS Foundation Trust**  
Heath Rd  
Ipswich  
United Kingdom  
IP4 5PD

**Study participating centre**  
**Basingstoke and North Hampshire Hospital**  
Hampshire Hospitals NHS Foundation Trust  
Aldermaston Road  
Basingstoke  
United Kingdom  
RG24 9NA

**Study participating centre**  
**Queens Hospital**  
Burton Upon Trent  
United Kingdom  
DE13 0RB

## **Sponsor information**

**Organisation**  
Sheffield Teaching Hospitals NHS Foundation Trust

**ROR**  
<https://ror.org/018hjpz25>

## **Funder(s)**

**Funder type**  
Government

**Funder Name**

## Results and Publications

### Individual participant data (IPD) sharing plan

The data sharing plans for the current study are unknown and will be made available at a later date.

### IPD sharing plan summary

Data sharing statement to be made available at a later date

### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Results article</a>		24/07/2024	25/07/2024	Yes	No
<a href="#">Results article</a>	Decisional regret	25/09/2024	26/09/2024	Yes	No
<a href="#">Basic results</a>		21/03/2024	28/03/2024	No	No
<a href="#">HRA research summary</a>			28/06/2023	No	No
<a href="#">Study website</a>	Study website	11/11/2025	11/11/2025	No	Yes