

Proper understanding of recurrent stress urinary incontinence treatment in women

Submission date 17/12/2019	Recruitment status No longer recruiting	<input type="checkbox"/> Prospectively registered
		<input checked="" type="checkbox"/> Protocol
Registration date 09/01/2020	Overall study status Stopped	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
Last Edited 11/09/2025	Condition category Urological and Genital Diseases	<input type="checkbox"/> Individual participant data
		<input type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

Urinary leakage with physical activity is called stress urinary incontinence (SUI), and it affects about a quarter of women after pregnancy. Until recently, the most common treatment was a surgical operation which helps to support the tube which takes urine from the bladder to the outside (urethra), called midurethral (mesh) tape. Unfortunately, symptoms can come back after treatment, this is called recurrent SUI. In some cases, symptoms may never have gone away, this is called persistent SUI. Current treatment options for recurrent or persistent SUI include:

1. Injections into the urethra to help it to seal when leaks might happen called endoscopic bulking injections. The injections are done from a tube outside the body.

2. Surgical operations include:

- A medical mesh tape is placed in the vagina to support the urethra (midurethral tape)
- A strip of the patient's own tissue (taken from the tummy area) is used to support the urethra (autologous fascial sling)
- Stitches are used to lift the vagina so that it supports the urethra (colposuspension)
- An implant device is placed around the urethra to gently squeeze it and prevent leaking (artificial urinary sphincter)

It is not known which of these treatments is best for women who have already had an operation or injections for SUI. The aim of this study is to find out whether surgical operations or endoscopic bulking injections are better for treating recurrent or persistent SUI.

Who can participate?

Adult (18 years or older) women with recurrent or persistent SUI who have already had an operation or bulking injection for it

What does the study involve?

250 women are recruited to the study. Equal numbers of women join an endoscopic bulking injection group or a surgical operation group. Which group women join will be decided by chance (in a process called randomisation). Women in the surgical operation group decide which operation to have with their doctor. Women receive their treatment and aftercare at hospital as they would during normal NHS care and are asked to complete a questionnaire booklet at the start of the study and again 6 months, 1, 2 and 3 years later. The questionnaires cover general health, urinary symptoms and the effect of those symptoms on everyday life and sex life. The

researchers audio-record consultations where the study is discussed with women and interview some women to see how research is explained and understand how women manage after their treatment.

What are the possible benefits and risks of participating?

Some people enjoy being part of research studies because of the close contact with research staff and the opportunity to share their opinions and experiences of their condition and treatments. Women will be offered a £10 voucher for completing their questionnaire at 1 year and another £10 voucher for completing their questionnaire at 3 years. There is no additional risk to normal NHS practice of the endoscopic bulking injections or surgical operations, and neither are new or experimental. Women taking part will have the same risks as anyone having treatment for recurrent SUI. This includes the possibility that symptoms may not improve as much as women would like. The risks and benefits of each treatment will be explained by the doctors, and women will be provided with relevant hospital leaflets.

Where is the study run from?

This study is sponsored by North Bristol NHS Trust. The Bristol Randomised Trials Collaboration (as part of the Bristol Trials Centre) at the University of Bristol is responsible for managing the study. The researchers aim to run the study in 20 NHS hospitals across the UK.

When is the study starting and how long is it expected to run for?

April 2019 to October 2022

Who is funding the study?

National Institute of Health Research (NIHR) Health Technology Assessment (HTA) Programme (UK)

Who is the main contact?

Dr Caroline Pope
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Contact information

Type(s)

Public

Contact name

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Additional identifiers

Clinical Trials Information System (CTIS)
Nil known

Integrated Research Application System (IRAS)
257547

Protocol serial number
Sponsor reference: #4404, HTA 17/95/03, IRAS ID 257547

Study information

Scientific Title
Proper Understanding of Recurrent Stress Urinary Incontinence Treatment in women (PURSUIT): a randomised controlled trial of endoscopic and surgical treatment

Acronym
PURSUIT

Study objectives
To determine whether surgical treatment is superior to endoscopic bulking injections in terms of symptom severity at 1-year after randomisation, in women with recurrent SUI.

Ethics approval required
Old ethics approval format

Ethics approval(s)
Approved 19/12/2019, South West - Frenchay Research Ethics Committee (Level 3, Block B, Whitefriars, Lewins Mead, Bristol, BS1 2NT, UK; Tel: +44 (0)207 1048 045; Email: nrescommittee.southwest-frenchay@nhs.net), ref: 19/SW/0209

Study design
Two-arm multi-centre interventional randomised controlled trial

Primary study design
Interventional

Study type(s)
Treatment

Health condition(s) or problem(s) studied
Recurrent or persistent Stress Urinary Incontinence (SUI)

Interventions
Participants will be randomised on a 1:1 basis using an online randomisation system or automated telephone system:

Arm 1 - endoscopic (urethral) bulking injections

Arm 2 - surgical procedure (colposuspension or autologous urethral sling or midurethral tape or artificial urinary sphincter (AUS)); women in the surgical operation group will decide which operation to have with their doctor

Women will receive their treatment and aftercare at hospital, as they would during normal NHS care and will be asked to complete a questionnaire booklet at the start of the study and again 6 months, 1, 2 and 3 years later. The questionnaires cover general health, urinary symptoms and the effect of those symptoms on everyday life and sex life. The researchers will audio-record consultations where the PURSUIT study is discussed with women and interview some women to see how research is explained and understand how women manage after their treatment.

Intervention Type

Procedure/Surgery

Primary outcome(s)

Patient-reported outcome measure (PROM) of continence using the International Consultation on Incontinence Questionnaire - Urinary Incontinence - Short Form (ICIQ-UI-SF) at 1 year after randomisation

Key secondary outcome(s)

1. Clinical subjective measure of continence (longer term) using the ICIQ-UI-SF questionnaire at 6 months, 2 and 3 years post randomisation
2. Improvement of symptoms measured using the Patient Global Impression of Improvement (PGI-I) questionnaire at 1, 2 and 3 years post randomisation
3. Procedure/operative assessment measures: assessment of procedure/operation time, estimated blood loss, hospital stay, and return to normal activity, measured at time of intervention and at 6 months post-intervention
4. Incontinence and sexual function assessed using the Pelvic Organ Prolapse/Incontinence Sexual Questionnaire (PISQ-IR) at 1, 2 and 3 years post randomisation
5. Adverse events: evaluation of treatment and retreatment, adverse events of each intervention at intervention, 6 months post intervention, and 6 months, 1, 2 and 3 years post randomisation
6. Cost-effectiveness from an NHS and societal perspective in terms of Quality-Adjusted Life Years (QALYs) and ICIQ-UI-SF at 1 year, and from a secondary care NHS perspective in terms of QALYs at 3 years. EQ-5D-5L (used to calculate QALYs) questionnaire at 6 months, 1, 2 and 3 years post-randomisation. Secondary care resource use from Trust electronic systems (or Hospital Episode Statistics) at 1 and 3 years post-randomisation. Community-based and patient resource use questionnaire at 6 months and 1 year post-randomisation
7. Patient experiences of the intervention, assessed using qualitative interviews with patients at 6 months, 1 year and 3 years post-intervention
8. Clinician views of the intervention, assessed using qualitative interviews with clinicians around baseline

Completion date

04/10/2022

Reason abandoned (if study stopped)

Lack of funding/sponsorship

Eligibility

Key inclusion criteria

1. Adult women (≥ 18 -years) with bothersome Stress Urinary Incontinence (SUI) symptoms after primary SUI surgery (including bulking injections)
2. Urodynamics to confirm recurrent or persistent SUI
3. Patient willing to consider interventional therapy
4. Patient willing to be randomised and willing to give consent

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Adult

Lower age limit

18 years

Sex

Female

Total final enrolment

23

Key exclusion criteria

1. Predominant urgency incontinence
2. Pelvic organ prolapse (POP) more than or equal to stage II
3. Relevant neurological disease, disease, such as a stroke, multiple sclerosis, Parkinson's disease, or spina bifida (diabetes mellitus is not an exclusion criterion unless it is causing diabetic neuropathy)
4. Being treated for gynaecological or bladder cancer
5. Unresolved mesh exposure from previous midurethral tape (MUT)
6. Current pregnancy
7. Urethral diverticulum
8. Recent pelvic surgery (e.g. POP repair, stress incontinence surgery, and hysterectomy within the last 6-months)
9. Participation in another study that might influence results or increase patient burden
10. Unable to give informed consent/complete assessments
11. Previous artificial urinary sphincter (AUS) surgery

Date of first enrolment

10/12/2019

Date of final enrolment

11/07/2022

Locations

Countries of recruitment

United Kingdom

England

Scotland

Study participating centre**North Bristol NHS Trust**

Southmead Hospital

Southmead Road

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Bristol

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BS10 5NB

Study participating centre**NHS Ayrshire and Arran**

PO Box 13

Boswell House

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Study participating centre**University College London Hospitals NHS Foundation Trust**

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NW1 2PG

Study participating centre**Birmingham Women's and Children's NHS Foundation Trust**

Steelhouse Lane

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B4 6NH

Study participating centre

Sheffield Teaching Hospitals NHS Foundation Trust

Northern General Hospital
Herries Road
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United Kingdom
S5 7AU

Study participating centre

Cambridge University Hospitals NHS Foundation Trust

Hills Road
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United Kingdom
CB2 0QQ

Study participating centre

South Tees Hospitals NHS Foundation Trust

The James Cook University Hospital
Marton Road
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United Kingdom
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Study participating centre

Royal Cornwall Hospitals NHS Trust

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TR1 3LJ

Study participating centre

East Lancashire Hospitals NHS Trust

Royal Blackburn Hospital
Haslingden Road
Blackburn
United Kingdom
BB2 3HH

Study participating centre

Leeds Teaching Hospitals NHS Trust

St. James's University Hospital

Beckett Street
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United Kingdom
LS9 7TF

Study participating centre
Stockport NHS Foundation Trust
Stepping Hill Hospital
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SK2 7JE

Study participating centre
Northern Care Alliance NHS Foundation Trust
Salford Royal
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M6 8HD

Study participating centre
Liverpool Women's NHS Foundation Trust
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L8 7SS

Study participating centre
Bedfordshire Hospitals NHS Foundation Trust
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Study participating centre
Mid and South Essex NHS Foundation Trust
Prittlewell Chase

Westcliff-on-sea
United Kingdom
SS0 0RY

Sponsor information

Organisation

North Bristol NHS Trust

Funder(s)

Funder type

Government

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

Current IPD sharing plan as of 11/09/2025:

Anonymous study data will be kept on a secure restricted-access University of Bristol server. Requests for access to the anonymised data set should be made via the Bristol Trials Centre (BTC, btc-mailbox@bristol.ac.uk) and Chief Investigator (CI, marcus.drake@imperial.ac.uk). Requests must be via a written confidentiality and data sharing agreement (DSA). The DSA should cover limitations of use, transfer to third parties, data storage and acknowledgements. The person applying for use of the data will be scrutinised for appropriate eligibility by the BTC /CI. The approved Participant Consent Form for the study includes the clause "I understand that the information collected about me will be used to support other research in the future and may be shared anonymously with other researchers."

Previous IPD sharing plan:

Anonymous study data will be kept securely on the University of Bristol Research Data Storage Facility (RDSF, <https://www.bristol.ac.uk/acrc/research-data-storage-facility/>). After the study is finished, requests for access to data should be made via the University of Bristol Research Data Repository (<https://data.bris.ac.uk/data/>). Requests must be via a written confidentiality and data sharing agreement (DSA) which will be confirmed/approved by the Chief Investigator. The DSA should cover limitations of use, transfer to third parties, data storage and acknowledgements. The person applying for use of the data will be scrutinised for appropriate eligibility by the research team/CI. The approved Participant Consent Form for the study includes the clause "I understand that the information collected about me will be used to support other research in the future and may be shared anonymously with other researchers".

IPD sharing plan summary

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol article		03/08/2022	04/08/2022	Yes	No
HRA research summary			26/07/2023	No	No
Study website	Study website	11/11/2025	11/11/2025	No	Yes