

Participant information sheet for doctors and researchers

<u>Developing Core Outcomes sets in Female</u> <u>Idiopathic Chronic Pelvic Pain</u>

(Study Number: 277330)

We would like to invite you to take part in our research study. Participation is entirely voluntary. Before you decide whether to take part in the study, we would like you to understand why the research is being done and what it would involve for you.

Chief Investigator:

Mr. Stergios K Doumouchtsis Department of Obstetrics and Gynaecology Epsom Hospital Dorking Road

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Epsom and St Helier WHS
University Hospitals

What is the purpose of the study?

This study is to find out what you think the priorities in research for chronic pelvic pain should be. By

involving doctors and researchers we can ensure your views are represented in future research.

Why have I been chosen?

We are inviting all doctors and researchers with an interest in female chronic pelvic pain to

participate in this research study.

Do I have to take part?

It is up to you to whether or not you choose to take part. You can find further information about the

study on our webpage https://i-chorus.org/core-outcome-sets. If you choose to take part in the

study you will need to register your details on our webpage. By registering on our website, you have

agreed and provided your consent to be involved in the study. However, your participation is

voluntary and you can withdraw at any time.

What will happen to me if I take part?

Once you register on our webpage, you will be also asked for an email address. This is so we can

contact you with instructions to complete the survey. There will be two surveys which can be

completed online. These will take 30 to 60 minutes each.

You will be also invited to attend a virtual meeting at the end of our study. This is to discuss the

results of the study. This meeting will be for half a day.

What are the possible risks of taking part?

There are no risks associated with taking part in the study. However, to minimise inconvenience to

you, surveys can be completed online over two weeks. You do not need to attend the meeting in

person and as the meetings will be held virtually

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Mr Stergios K Doumouchtsis

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Epsom and St Helier MHS **University Hospitals**

What are the possible benefits of taking part?

There is no direct benefit to you from taking part in the study. However, findings from this study can

be used to improve the care of women with chronic pelvic pain.

Will I be informed of the results?

Yes, we will provide a short summary and write our reports in a way that no-one can work out that

you took part in the study.

Can I opt out after agreeing to take part in the study?

You can withdraw at any time without giving a reason but we will keep the information we have

already about you.

How will we use information about you?

We will need to use information about you for this project. This will include your email address,

name, age, gender and survey responses. People will use these details to do the research or check

your records to make sure that the research is being done properly. People who do not need to

know who you are will not see your name or contact details. Your data will have a code number

instead. We will keep all information about you safe and secure. Once we have finished the study,

we will keep personally identifiable data so we can check the results for 12 months. We need to

manage your records in specific ways for the research to be reliable. This means that we won't be

able to let you see or change the data we hold about you.

The Sponsor has contracted the services of University of Liverpool to deliver the survey. They survey

will collect personal information such as your name, age, gender and email address which will be

held by the University of Liverpool in a secure database. This data will be held for 3 months after the

study ends and then destroyed. If you wish to request your personal identifiable data is removed

from the University of Liverpool database please contact the Chief Investigator.

Where can you find out more about how your information is used?

You can find out more about how we use your information:

On www.hra.nhs.uk/information-about-patients/

by asking one of our research team: Vishalli Ghai, vishallighai@nhs.net

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What if something goes wrong?

Please speak to a member of the research team if you have if you have any concerns about this

study. We will do our best to answer your questions. If you are still unhappy and want to make a

formal complaint, you may do this through the independent Patient Advice and Liaison Service

(PALS). (Contact: 01372 735 243; Mon-Fri 0930 -1530).

What will happen to any feedback I give?

We value any feedback on our study that will be used to further improve our research process.

Who is organising this study?

Epsom and St. Helier University Hospitals NHS Trust are sponsoring this study.

As part of the study, we have established an international steering group. This group has doctors, nurses and researchers as well as women who have chronic pelvic pain. The purpose of this steering

group is to help develop and guide the study.

Who is funding this study?

This project has been supported by a financial grant from Bayer Plc. Bayer Plc has had no

involvement whatsoever in the development or implementation of the project.

Who has reviewed the study?

This study proposal was submitted online to Health Research Authority and has been reviewed by a

Research Ethics committee West Midlands - Black Country Research Ethics Committee.

Contact for Further Information

For any further information please contact:

Mr. Stergios K Doumouchtsis, Epsom Hospital, Dorking Road, Epsom, KT18 7EG

Email address: s.doumouchtsis@nhs.net

To register your interest or for further information please visit [xxx] website

Thank you for considering taking part in this study.

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