**PARTICIPANT INFORMATION SHEET – PATIENTS AND CARERS**

Study Title: Evaluating video and hybrid group consultations

## We'd like to invite you to take part in our research study. Before you decide, it is important that you understand why the research is being done and what it would involve for you. Please take time to read this information and discuss it with others if you wish. *If there is anything that is not clear, please ask us. You can also find further information about how researchers use information from patients at;* [www.hra.nhs.uk/patientdataandresearch](http://www.hra.nhs.uk/patientdataandresearch)

# What is the purpose of the study?

Some GP practices in the UK invite patients to take part in clinical consultations as a group. These group sessions usually involve 6-8 patients with the same condition or similar health needs. Patients discuss test results, any medication changes and/or other health concerns with a clinician, and also provide support to each other in a group setting. In response to the Covid-19 pandemic, some of these group sessions are now delivered online on video, or involve some people joining online and others in-person in the same ‘hybrid’ session.

There are benefits and limitations to this approach of receiving care and we would like to better understand your views and experiences of video and hybrid group consultations. The aim of the study is to understand what it has been like for different people being involved in group consultations using online and hybrid formats. We want to understand what has worked well, not so well, and if or how the service could be improved. The study will also examine if there are differences between those attending group consultations and those attending individual appointments, in terms of how they feel about their health, how much they need to use health services and how satisfied they are with the service. Finally, we will look at what the costs are for delivering group consultations.

# Why have I been invited to take part?

You have been invited to take part in this research study, because you (or the person you care for) have participated in a video or hybrid group consultation, or have declined an invitation to participate in a video or hybrid group consultation. All participants should be over 18 years old. We plan to recruit 500 patients and carers to this study in total. We are also involving 20-30 staff, and 10-17 commissioning and policy stakeholders.

# Do I have to take part?

# No, taking part in this study is voluntary and you can withdraw at any time without giving a reason if you later change your mind.

# What will happen to me if I decide to take part?

If you are happy to participate in this study, you will be asked to provide consent. A copy of the record of consent will be given to you or sent by email for you to keep. Depending on what you prefer, you may be involved in one or more aspects of the study:

1. If you are taking part in a group consultation, a researcher may observe the group consultation, and they may ask your group some questions at the end of the session to understand what has worked well and what hasn’t.
2. A researcher may interview you separately about your experiences taking part in group consultations or your views about this way of receiving care, if you have declined to attend group sessions. The interview will take place by phone, online or in-person, depending on your preferences and Covid-19 restrictions. If you are willing, we may follow up with a second interview 12 months later to understand how things have changed.
3. You may be asked to attend a focus group discussion with 6-8 patients/carers in total, where a facilitator will invite you to discuss your views about group sessions in general practice.
4. You may be invited to fill in a one-off survey questionnaire at the end of one of your group consultations (those attending group consultations for the first time may be asked to fill in this questionnaire again if they attend another group consultation during the course of the project). We would like to collect different types of information to find out more about you, your health, your confidence at managing any chronic conditions, and your experience with general practice.
5. We would like to obtain data from your health records so that we can understand more about your health and usage of health services (over a period of 12 months from involvement).
6. You may also be asked to complete a health-related quality of life and satisfaction questionnaire (called Measures of Health Benefits survey) after your menopause consultation. This will help us to understand in greater detail about your experiences of your health condition(s) and your experiences about the consultation process itself.

With your consent, interviews and focus groups will be recorded via audio (e.g. over the phone, online or face-to-face). Group consultations will be recorded via audio when observations take place in-person and/or on video (e.g. using built-in recording functionality and/or video camera). Alternatively the researcher can take verbatim notes.

In some cases we will ask a small number of selected patients who participate in video/hybrid group consultations remotely, if they are happy for us to join them (e.g. at home) and observe how the group consultation works from their perspective. With consent, we will place a small video camera near your computer screen to capture your interaction with the clinician and rest of patients. If you want, the researcher will leave the room during your consultation.

**What if I need help from a health or support worker?**

You can ask your support worker, a family member or someone you trust to help you.

You can let them know about this information sheet and confirm with them if you want to take part. If you agree, your support worker will pass on your contact details to the researcher. They will contact you and will find a day or time to talk that will suit you.

# What should I consider?

# The main thing to consider is whether you are comfortable with sharing your experiences and views on primary care services. This could raise some potentially sensitive subjects around your health or care experiences.

# Are there any possible disadvantages or risks from taking part?

There are minimal risks in this study other than time commitment. If you agree to be interviewed, this will take up 30-45 minutes of your time, and if you agree to take part in a focus group, this will last up 60-90 minutes. One off (experience) survey questionnaires will take between 10-15 minutes. If you agree to take the health-related quality of life and satisfaction survey it will take a further 10-15 mins to complete. The research team will be able to see and hear your group consultation with other patients. If there are specific topics that you do not wish to be included, you can inform the researcher. Recordings will be de-identified using video-editing software according to established guidelines. If at any point during the consultation or interview, you feel any discomfort or distress, please inform your clinician or the researcher and the recording will be paused immediately. We will check if you are okay, and if you are happy to proceed.

# What are the possible benefits of taking part?

# Whilst we cannot guarantee any direct benefit to you, our aim is to use the research results to help improve the NHS. Learning from this study could potentially lead to improved organisation of remote group-based service delivery in general practice, better patient care and self-management.

# Will I be reimbursed for taking part?

# There will be no reimbursement other than compensation for travel expenses if you need to travel to take part in the study beyond your usual care.

# Will my taking part in the study be kept confidential?

Yes. People who do not need to know who you are will not be able to see your name or contact details. Your data will have a code number instead. We will keep a separate paper record in a locked cabinet of your real name and corresponding code number.

If anything is said that raises a concern of harm to self or others, or suggests poor practice, we have safeguarding obligations to follow this up, which may mean discussing with others.

We will write our reports in a way that no-one can work out that you took part in the study. The recordings will be de-identified using video-editing software according to established guidelines, in which we remove any spoken identifiers (e.g. names) and apply a visual filter to make you unrecognisable and remove any names from the audio track.

Responsible members of the University of Oxford and the relevant GP practice may be given access to data for monitoring and/or audit of the study to ensure that the research being done properly.

### **What will happen to my data?**

UK Data protection regulation requires that we state the legal basis for processing information about you. In the case of research, this is ‘a task in the public interest.’ The University of Oxford, based in the United Kingdom is the sponsor for this study and the data controller, and is responsible for looking after your information and using it properly.

We will be using information from you and your medical records to do this research and will use the minimum personally-identifiable information possible. People will use this information to do the research or to check your records to make sure that the research is being done properly. We will keep all information about you safe and secure on a non-networked computer at the Universities of Oxford, York and Exeter.

We will keep identifiable information about you including your name and email or telephone number for 12 months after the study has finished, so as to contact you about the research study and feed back results should you so wish. All interview recordings will be destroyed after transcripts have been checked for accuracy.

However, research documents with personal information, such as consent forms, will be held securely at the University of Oxford for 15 years after the end of the study. We will keep some of the data so we can check the results.

Some of your information will be sent to our research partners at the University of York and University of Exeter, and a professional transcription service (i.e. they write out the text of the whole interview). They must follow our rules about keeping your information safe by signing a confidentiality agreement. Identifiable information about you will be removed from audio files before sharing with them. Transcribers will destroy their copy of the recording once they have completed transcription.

All data from recorded interviews will be de-identified when they are transcribed. This means that identifiable information about participants will be removed from transcripts (i.e. the text from the interview is written out) or other notes researchers may take.

Data, including recordings of the video and group consultations, will be handled in line with UK General Data Protection Regulation (GDPR) and the Data Protection Act (2018).

UK Data protection regulation provides you with control over your personal data and how it is used. When you agree to your information being used in research, however, some of those rights may be limited in order for the research to be reliable and accurate. Further information about your rights with respect to your personal data is available at

<https://compliance.web.ox.ac.uk/individual-rights>

You can find out more about how your information is used by asking one of the research team

or by emailing the Chief Investigator (sara.shaw@phc.ox.ac.uk )

You can find general information about how patient information is used in researchat [www.hra.nhs.uk/information-about-patients/](https://www.hra.nhs.uk/information-about-patients/)

# What will happen to my data at the end of the study?

# The members of our research team will analyse the data and write some papers and reports, including a summary for the general public. Our findings will be published and available through journal publications. You will not be identified from any report or publication placed in the public domain. We may wish to use quotes from your interview or our notes in a conference presentation, academic article or teaching session, but you would not be identifiable and do not have to agree to this. If you agree for us to use video clips or stills from your group consultation in presentations and publications, these will be pseudonymised and in pixelated format (we will apply a visual filter, remove names, and mask voices). If you are happy for this to happen, please indicate during the consent process.

**Can I change my mind about participating?**

You can stop at any time, without giving a reason and without penalty, by telling the researchers of your decision (using the contact details at the end of this information sheet). Your participation is voluntary and even if you originally said yes, you may change your mind at a later stage (any time before the project end date in November 2024). The researchers will ask for your permission to keep data already collected but you are free to refuse this. Please note information from your consultations is also recorded by the GP surgery as part of standard clinical care and the study team has no control over how this information is processed.

A member of our study team may wish to record a reason about why you have withdrawn for our record keeping, but you do not have to provide one.

# What if there is a problem?

The University of Oxford, as Sponsor, has appropriate insurance in place in the unlikely event that you suffer any harm as a direct consequence of your participation in this study.

If you wish to complain about any aspect of the way in which you have been approached or treated, or how your information is handled during the course of this study, you should contact the Chief Investigator Prof. Sara Shaw by email ([sara.shaw@phc.ox.ac.uk](mailto:sara.shaw@phc.ox.ac.uk)) or you may contact the University of Oxford Research Governance Ethics and Assurance (RGEA) office on 01865 616480, or the director of RGEA, email [rgea.complaints@admin.ox.ac.uk](mailto:ctrg@admin.ox.ac.uk).

The Patient Advisory Liaison Service (PALS) is a confidential NHS service that can provide you with support for any complaints or queries you may have regarding the care you receive as an NHS patient. PALS is unable to provide information about this research study.

You can contact the local NHS Patient Advice and Liaison Service (PALS), at 01865 221473 or here: <https://www.nhs.uk/service-search/other-services/Patient-advice-and-liaison-services-(PALS)/LocationSearch/363>

# How have patients and the public been involved in this study?

Patient and public contributors helped design the research study and data collection tools.

# Who is organising and funding the study?

# The project is led by Drs Chrysanthi Papoutsi and Sara Shaw at the Department of Primary Care Health Sciences and funded by the National Institute for Health Research through the Health Services and Delivery Research (HS&DR) Programme (with small add-on funding provided by the National Institute for Health Research School of Primary Care Research).

# Who has reviewed the study?

All research in the NHS is looked at by an independent group of people, called a Research Ethics Committee, to protect participants’ interests. This study has been reviewed and given favourable opinion by the London - Hampstead Research Ethics Committee.

# Further information and contact details:

If you would like to discuss the research with someone beforehand (or if you have questions afterwards), please email the Chief Investigator:

Prof Sara Shaw, Nuffield Department of Primary Care Health Sciences, University of Oxford, Radcliffe Primary Care Building, Woodstock Rd, Oxford OX2 6GG

E: sara.shaw@phc.ox.ac.uk

*Thank you for considering taking part.*