

**Feasibility and ACceptability of a mobile Technology intervention to Support [FACTS]  
Post Abortion Care in British Columbia Study**

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## Study information Sheet – User Testing

### What is the FACTS study?

Individuals who have undergone an abortion have expressed there is a lack of follow-up care, especially those who live in rural and remote areas. We are creating a web-based platform to support follow-up care after an abortion procedure. This platform will include features relating to post-procedural care, psychosocial support, contraception decision-making and other resources relating to sexual health.

### Who can participate in this study?

- Individuals who have undergone an abortion procedure.
- Individuals who are fluent in English.

### What will your participation involve?

We are inviting you to participate in a usability testing session where you will test the prototype and provide feedback and recommendations as an expert end-user on an anonymous electronic evaluation questionnaire. We want to find out how useful and relevant our support tool is for post-abortion care, and get your advice on the content, design, and implementation of the intervention.

### How much time is required to participate?

Your total participation time will be 90 minutes, which will give you time to look through the different features of the website, and answer the evaluation form.

### What are the potential benefits of participating?

There may or may not be direct benefits to you from taking part in user testing of our prototype. We hope that the feedback you provide will further refine our platform to be the most useful and beneficial tool for individuals who undergo an abortion procedure in the future.

### **What happens if I decide to withdraw my consent to participate?**

You may withdraw from this user testing at any time without giving reasons. However, because the data are completely anonymous, any answers you provide prior to stopping will remain in the dataset.

### **Will I be paid to participate in this study?**

Participants will receive a \$10 e-gift card for participating.

### **How will my taking part in this study be kept confidential?**

The website evaluation form is anonymous, and your responses will not be linked to your personal identity. On completion of the survey the research coordinator will send you the e-gift card honorarium for your participation to your email address.

### **Who do I contact if I have questions about the study during my participation?**

If you have any questions or desire further information about this study before or during participation, or if you experience any adverse effects, you can contact Dr. Roopan Gill at [Roopan.Gill@cw.bc.ca](mailto:Roopan.Gill@cw.bc.ca) or the Study Coordinator, Ciana Maher, 604-875-2424 ext. 4978, [Ciana.Maher@cw.bc.ca](mailto:Ciana.Maher@cw.bc.ca).

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Participant Consent

My participation in this study means:

- I have had enough time to think about the information provided.
- I have been able to ask for advice if needed.
- I have been able to ask questions and have had satisfactory responses to my questions.
- I understand that all of the information collected will be kept confidential and that the results will only be used for scientific purposes.
- I understand that my participation in this study is voluntary.
- I understand that I am completely free at any time to refuse to participate or to withdraw from this study at any time, and that this will not change the quality of care that I receive.
- I understand that I am not waiving any of my legal rights as a result of participating in this study.
- I understand that there is no guarantee that this study will provide any benefits to me.
- I have read and understood the information in this consent statement

By completing this survey, I consent to participate in this study.