BMJ Open Tablet computers for implementing NICE antenatal mental health guidelines: protocol of a feasibility study

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ABSTRACT

Introduction: Depression is one of the most common mental health disorders that may affect women during pregnancy. The prompt identification of this disorder, and the provision of treatment, may help to reduce the likelihood of post-partum depression, prevent severe forms of the disease, and reduce its intergenerational impact. Despite women's repeated encounters with health services throughout their antenatal care, depression often goes undiagnosed. This is one area where mobile health could prove useful. We will assess the feasibility of using tablets to incorporate depression screening into antenatal pathways. We will also assess if survey layout could affect the quality of the data collected through these devices.

Methods and analysis: We will test the feasibility of using iPad Airs for the administration of the Whoolev questions and the Edinburgh Postnatal Depression Scale (EPDS) to pregnant women attending antenatal clinics in England. We will assess the impact of survey layout on the quality of the responses given to these screening scales using a parallel, randomised controlled study design. We will calculate the positive predictive value, the negative predictive value and the false omission rate of the Whooley questions in comparison with the EPDS. We will calculate differences in data equivalence, time needed to complete the surveys, break-off rates, data completeness and requests for help between the 2 experimental groups: using all questions in one screen and navigation by vertical scrolling, or a single question per screen and navigation by multiple pages. Ethics and dissemination: This study has been

approved by the National Research Ethics Service Committee South East Coast—Surrey. Our findings will be disseminated through academic peer-reviewed publications, conferences and discussion with peers.

INTRODUCTION

Maternal mental health is a key public health priority due to the impact it can have on a woman's well-being and on her child's emotional, behavioural, cognitive and social development.^{1–5} Perinatal depression is one of the most common psychiatric disorders that can affect up to 15% of women during

Strengths and limitations of this study

- This study addresses an important area of unmet clinical need, which has a considerable impact on healthcare systems and society in general.
- This study will attempt to identify implementation issues (arising from the introduction of an mHealth intervention into a clinical process) that could impair the ability of already busy healthcare professionals to perform their clinical duties. This is in contrast to some existing studies, where mHealth interventions are deployed in highly controlled environments that do not capture the realities of clinical settings.
- This study will explore the effect that survey presentation could have on the quality of the data collected (when using a validated scale). This is in contrast to other studies in the medical literature (that assess mHealth interventions), which have not explored how these additional factors could affect data quality.
- This study will explore mHealth as a delivery mode for a depression screening validated scale. However, it will not evaluate the potential of mHealth as a delivery mode for evidence-based interventions.
- Our findings might only be generalisable to devices with similar technical specifications that are being used in similar clinical settings.

pregnancy or within 1 year of giving birth. The direct and indirect financial costs of perinatal depression to the health system, together with its direct impact on the mother and her child, account for the majority of the overall costs to society of perinatal mental disorders.⁵ In the UK, these costs have been estimated at £8.1 billion for each 1-year cohort of birth (with the average costs of one case of perinatal depression estimated at £74 k), 72% of which could be attributed to the adverse effects experienced by the child.² For this reason, the early identification of women who might be at risk of

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developing or have perinatal depression should be considered a priority.⁶

Women appear to be at an increased risk of depression 3–6 months after childbirth,^{1 4} although a recent longitudinal study found that symptoms of depression may be more prevalent at 4 years post-partum.⁷ Several factors are known to increase the likelihood of developing postpartum depression.^{1 6 8} Depression during pregnancy, however, has been recognised as one of the most important risk factors and one that is susceptible to early intervention. The point prevalence of depression during the first, second and third trimesters of pregnancy has been estimated at 7.4%, 12.8% and 12%, respectively.⁹ The timely identification of depression during these stages, and the provision of appropriate treatment, can reduce the likelihood of developing post-partum depression, prevent more severe forms of this disorder, and improve a woman's general health status.^{4 5} Treating depression during pregnancy may also help reduce the intergenerational impact of this condition.

Nonetheless, depression during pregnancy often goes undiagnosed and its routine screening is still debated.¹⁰ Validated screening tools, such as the Edinburgh Postnatal Depression Scale (EPDS), can facilitate the early identification of depressive symptoms in pregnant women.¹¹ However, cost-effectiveness is often cited as one of the key reasons against the routine screening for this condition.¹² This is an area where smartphones and tablet computers could demonstrate their value in a field of study that is known as mobile health (mHealth). Thanks to their advanced computational and networking capabilities, and overall popularity (in the first quarter of 2015, it was reported that approximately 66% of adults in the UK owned a smartphone),¹³ these devices could facilitate depression screening and monitoring of symptoms¹⁴ while reducing costs and administrative burden.

Researchers from the School of Social Work at the and staff University of Illinois members at Champaign-Urbanara Public Health District will be testing the use of tablet computers to implement depression screening in maternal clinics.¹⁵ They will be doing so during antenatal consultations. Mobile devices could be used to screen for depression before a consultation, thus releasing some of the consultation time. For this reason, we need to establish the feasibility of using these devices to do so.

One aspect of delivering depression screening through mobile devices is the impact that this technology may have on data quality. This is to ensure that the data accurately reflect underlying clinical changes. Our recent Cochrane review¹⁶ suggests that collecting patient self-reported data through apps is equivalent to collecting data through alternative delivery modes (ie, paper, short messaging service, laptops, plastic rulers and personal digital assistants). This review also highlighted that equivalence, and other data quality dimensions, could be affected by factors other than the delivery mode.¹⁶ One such factor is survey presentation. Unlike the survey

methodology literature, however,¹⁷¹⁸ the effect of this factor on validated clinical tools has not been systematically explored in the medical literature.

In this study, we will assess the feasibility of implementing the recommendations from The National Institute for Health and Care Excellence (NICE)¹⁹ for recognising depression during pregnancy. We will conduct our study in the waiting area of general practices, community midwiferies and hospitals using iPad Air tablets. We will evaluate the statistical properties of one of the validated instruments recommended by this guideline. We will also assess if survey layout (as a component of survey presentation) affects the quality of the responses collected through the screening instruments under evaluation.

METHODS AND ANALYSIS Study design

We will assess the feasibility of using iPad Air tablets to administer a brief survey asking for personal information, the Whooley questions and the EPDS (see online supplementary appendix 1) to pregnant women attending antenatal clinics across a number of primary and secondary care National Health Service (NHS) centres in England. iPad Air tablets have a 9.7-inch (diagonal) retina display, with a 2048-by-1536 resolution at 264 pixels per inch. Study data will be collected and managed using Snap survey tools.²⁰ Snap Mobile app V.4.0.30 (or later, if a new version becomes available) will be running on iOS V.9.0.2 (or later, if a new version becomes available).

In order to evaluate the effect of survey layout on data quality, we will use a parallel, randomised controlled study design. Pregnant women consenting to take part in this study will be randomly assigned to complete (1) an app version of the Whooley questions and the EPDS in which all the questions are presented on a single screen (app screening—scrolling layout), causing participants to scroll vertically on the device; or (2) an app version of the Whooley questions and the EPDS in which only one question per page is presented at any given time (app screening—paging layout), causing participants to navigate through multiple pages.

Pilot phase

We piloted both versions of the surveys used in this study (paging layout and scrolling layout) with four staff members at the Department of Primary Care and Public Health at Imperial College London. This phase allowed us to identify (and rectify) the following issues:

- ► Font size;
- Consistency of response formats (in particular, observed differences between yes-no questions and multiple choice questions with a single answer);
- ▶ Framing of the survey results.

Sample selection and recruitment

We will select our sample of participants from pregnant women attending antenatal clinics in general practices, midwifery services and hospitals ,which are part of the NHS in England. Each potential participant will be assessed by clinical staff against our inclusion and exclusion criteria (table 1).

On the day in the antenatal clinic, potential participants will be first approached in the waiting room of the participating centres by a member of the clinical care team. If a potential participant expresses her interest in this study, she will be introduced to one of our researchers who will explain the details of the study to her. Potential participants will be provided with a patient information sheet. The researcher will emphasise that the potential participant can ask any questions regarding the study. If a potential participant agrees to take part in this study, she will be asked to complete a consent form, followed by the study procedures (ie, personal demographic survey, Whooley questions, and Edinburgh Postnatal Depression Scale). Otherwise, if she is unsure about participation, she will have at least 24 h to decide. In this case, she would only take part in the study if she comes into contact with the research team during a future antenatal clinic.

Refusal to take part in this study will not have an impact on a woman's legal rights, medical care or on the relationship with her care providers. We will obtain written, informed consent from those women who, after receiving all the relevant study information and asking study-related questions, still wish to take part in this study.

Interventions to be measured Surveys

Non-validated, personal demographic survey

We will administer an 11-question survey to collect information about the woman's age group, ethnic background, marital status, employment status, level of education, smartphone or tablet computer ownership, parity, and previous personal history of depression.

Inclusion criteria	Exclusion criteria	
Women who are 18 years old or older	Current diagnosis of mood (affective) disorders as specified in the ICD-10, Classification of Mental Health and Behavioural Disorders ²¹	
Pregnant and of any gestational age Any parity	Currently receiving treatment for mood (affective) disorders Recent personal history of mood (affective) disorders within the past 12 months	
Attending antenatal clinics in participating NHS sites	Not comfortable reading and writing in English	

ICD-10, 10th revision of the International Statistical Classification of Disease and Related Health Problems; NHS, National Health Service.

Whooley questions

The Whooley questions were developed as a case-finding instrument for depression in primary care.²² This twoquestion instrument assesses depressed mood and anhedonia that have been present during the past month. Respondents are required to answer *yes* or *no* to each question.

Edinburgh Postnatal Depression Scale

The EPDS is a 10-item self-administered survey that was developed to screen for perinatal depression in the community.²³ This instrument assesses feelings of guilt, sleep disturbance, reduced energy levels, anhedonia and suicidal ideation that have been present during the past 7 days. Each question is scored on a four-point scale ranging from 0 to 3 points. An overall score is calculated by adding the scores from each individual question. Overall scores between 10 and 12 points suggest increased risk for depression; scores of 13 points or more suggest that the diagnostic criteria for major depression disorder have probably been met.²⁴ In addition, special attention should be paid to item 10, as it deals with suicidal thoughts. On the basis of these scores, a clinician would be prompted to refer a woman to a mental health professional.

The EPDS is a valid and reliable tool for identifying women who are at risk of depression, both during pregnancy and post-partum. This instrument is also sensitive to changes in the severity of depression over time.²³ The EPDS can be reproduced without further permission provided that the original source of the scale is cited in each reproduced copy.

Screening for depression and effect of survey layout

Pregnant women taking part in our study will receive verbal instructions on how to complete the surveys. Participants will also be informed that we will communicate their answers to the Whooley questions and their scores on the EPDS to their clinicians, and that these results will be available for discussion during their antenatal appointment. Participants' results on the Whooley questions and on the EPDS will be communicated to their general practitioners.

The researcher will also explain to participants that they can ask for help at any point while completing the surveys, and that all questions should be directed to a member of the research team (not to the clinical care team). We will offer help with the use of the device. However, we will not offer help reading questions out loud to participants (as the Whooley questions and the EPDS are intended to be used as self-administered instruments).

Participants will be asked to complete all the surveys while waiting for their appointment in the waiting room of participating NHS sites. They will be randomised to one of two experimental manipulations of the survey layout.

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Scrolling layout

In this experimental manipulation, all the surveys will be presented on a single screen (figure 1). Therefore, participants will need to scroll vertically in order to answer all the questions. They will be able to scroll up and down as they wish and to modify the answers given to previous questions as long as they do it before submitting their answers. Once a participant has submitted her answers, a message acknowledging their participation will be displayed on the screen of the device and they will no longer be able to modify their answers. Once all the surveys have been completed, participants will be asked to return the iPad Air to a member of the research team. Validation procedures built into Snap surveys will ensure that participants are not able to submit unanswered questions.

Paging layout

In this experimental manipulation, participants will only see one question on the screen at any given time (figure 2). They will need to select and submit their answer to the current question before being able to move on to the next question. Participants will not be allowed to revisit already answered questions. Once a participant has submitted all her answers, a message acknowledging her participation will be displayed on the screen of the device. Once all the surveys have been completed, participants will be asked to return the iPad Air to a member of the research team. Validation procedures built into Snap surveys will ensure that participants are not able to submit unanswered questions.

Randomisation

We will utilise a block randomisation procedure to allocate participants to one of the two experimental arms. Random numbers will be generated using Stata V.13.0 (Stata. Stata Statistical Software[Program]. Version 13, 2013). Each consecutive number that is generated will be printed and put inside an opaque envelope, which will then be sealed. Envelopes will be numbered in an ascending sequence. Once a participant has provided informed consent, the researcher will take the relevant envelope, open it and use the number contained in it to allocate the participant to one of the two experimental groups. Researchers conducting participant recruitment will not be involved in this randomisation procedure in order to avoid recruitment bias.

Outcomes

Positive predictive value of the Whooley questions

We will calculate the number of pregnant women who answered *yes* to any of the Whooley questions AND who scored 10 points or higher on the EPDS, as a proportion of the total number of pregnant women who answered *yes* to any of the Whooley questions regardless of their EPDS scores.

Negative predictive value of the Whooley questions

We will calculate the number of pregnant women who answered *no* to both Whooley questions AND who scored nine points or less on the EPDS, as a proportion of the total number of pregnant women who answered *no* to both Whooley questions regardless of their EPDS scores.

False omission rate of the Whooley questions

We will calculate the number of pregnant women who answered *no* to both Whooley questions AND who scored 10 points or higher on the EPDS, as a proportion of the total number of pregnant women who answered *no* to both Whooley questions regardless of their EPDS scores.

Mean overall scores on the EPDS

For each experimental group, we will sum the individual scores on the EPDS and divide the total by the number of participants in the group. We will also calculate the mean overall scores on the EPDS for our total sample.

Break-off rates

We will calculate the proportion of participants who interrupt the survey before reaching the end. In such eventualities, we will document the reason for the break-off.

Time needed to complete surveys

We will measure the time in seconds between a participant starting to read the basic instructions on how to complete the survey and the participants receiving the message acknowledging their participation. We will also account for the time that participants spend experiencing problems, distractions or making requests for help or clarification.

Proportion of complete EPDS records

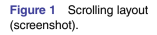
We will define a complete record as one in which no question was left unanswered. Leaving unanswered questions will not be possible due to the implementation of data validation procedures. Therefore, data completeness will be a by-product of break-off rates.

Proportion of participants requesting help

We will document the requests for help made by our participant. We will categorise participants into three groups: (1) one request for help; (2) between two and four requests for help; and (3) five or more requests for help. We will compute the proportion of participants falling within each category.

Proportion of each type of request for help

We will document the reasons for each request for help. This information will be captured through a post study procedures survey that researchers will have to complete. We will conduct a content analysis of these requests in order to categorise them according to their nature.



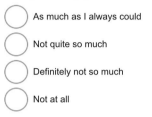
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Section 3: Edinburgh Postnatal Depression Scale

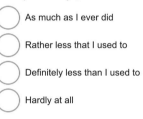
As you are pregnant, we would like to know how you are feeling.

Please check the answer that comes closest to how you have felt in the past 7 days, not just how you feel today.

Q3.1 In the past 7 days, I have been able to laugh and see the funny side of things:



Q3.2 In the past 7 days, I have looked forward with enjoyment to things:



Q3.3 In the past 7 days, I have blamed myself unnecessarily when things went wrong:



Q3.4 In the past 7 days, I have been anxious or worried for no good reason:

No, not at all

Sample size calculations

We powered our study according to the positive predictive value (PPV), negative predictive value (NPV) and false omission rate (FOR) of the Whooley questions. In order to calculate these outcomes, we would need at least 30 events (defined as pregnant women scoring 10 points or more on the EPDS). Assuming a prevalence of depression in pregnant women of 12%, we would need to recruit 250 participants in our study. We decided to recruit a total of 300 participants to account for any eventualities, such as dropouts.

Data analysis plan

Descriptive statistics

We will report the number of participants approached who did not meet our inclusion criteria, as well as the reason for exclusion. We will also report the number of participants who refused to take part in our study as a proportion of the total number of participants who were initially approached.

We will report the following information for each experimental group:

- Demographic characteristics (as captured by our non-validated survey);
- Proportion of participants answering *Yes* to any of the Whooley questions;
- Proportion of participants scoring at each interval of the EPDS: between 0 and 9 points; between 10 and 12 points; and 13 points or above;
- Proportion of participants scoring 2 or 3 points on question 10 of the EPDS;
- ▶ Mean time taken to complete the surveys;

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Q3.1	In the past 7 days, I have been able to laugh and see the funny side of things:
	As much as I always could
	Definitely not so much
	Not at all
	← Back Next →

- Proportion of participants making requests for help;
- Proportion of participants making each type of request;
- Mean time spent dealing with requests for help; and
- ▶ Break-off rates and reasons for break-off.

Inferential statistics *PPV*, *NPV and FOR*

We will calculate the PPV, NPV and FOR of the Whooley questions for the total sample of participants, and will report these values as percentages. We will also test for significant differences between the two experimental groups in these outcomes using a t test.

Data equivalence

We will determine data equivalence between the experimental groups (scrolling and paging) by testing for statistically significant differences in the mean overall scores on the EPDS. For this, we will conduct a t test. In addition, we will compare the proportion of women scoring (1) between 10 and 12 points, and (2) 13 points or more on the EPDS between the two experimental groups using a t test.

Time needed to complete surveys

We will compare the mean time needed by participants to complete the survey questionnaire between the experimental groups using a t test.

Data completeness

We will compare the proportion of complete records between the experimental groups using a t test.

Proportion of requests for help

We will compare the proportion of participants making requests for help between the experimental groups. For this, we will conduct a t test for each category in this end point: (1) one request; (2) between two and four requests; and (3) five or more requests.

Timeline

We expect recruitment of participants to take place between October 2015 and the end of February 2016.

CONCLUSION

We believe that this study addresses an important area of unmet clinical need which has a direct and indirect impact on health systems and society. It will also contribute to building the evidence base for mHealth, particularly when applied to mental health. Unlike other studies in the medical literature, this study will evaluate how survey layout could affect the quality of the responses collected using a validated scale. Evaluating how different factors could affect data quality becomes important when data are used to inform clinical and therapeutic decisions. In addition, this study will also evaluate some of the implementation issues that might arise when deploying an mHealth intervention, and that could affect its successful adoption.

The technical specifications of a device, such as screen size, could influence user interaction and thus the quality of responses. In addition, usage patterns (in terms of the type of interactions, duration and frequency of an interaction, and the type of activities that occur during an interaction) vary depending on the type of device and the setting in which it is being used. For these reasons, our findings might only be generalisable to iPad Air tablets (or devices with similar technical specifications) that are being used in clinical settings. Therefore, in a future study, we will explore issues associated with the at-home collection of mood data during pregnancy, using participants' own devices.

The findings from this feasibility study will inform a larger trial evaluating the integration of this intervention into routine antenatal pathways, and the potential impact on clinical outcomes.

ETHICS AND DISSEMINATION

This study has been reviewed and approved by the National Research Ethics Service (NRES) Committee South East Coast—Surrey on 9 July 2015 under the

Research Ethics Committee (REC) reference 15/LO/ 0977.

This study is being sponsored by Imperial College London under the reference number 15IC2687 and has been included in the UK Clinical Research Network (CRN) Study Portfolio under the UKCRNID number 19280.

Lastly, this study protocol has been registered in ClinicalTrials.gov under the identifier NCT02516982 (as required by the REC).

The findings of this study will be disseminated through academic peer-reviewed publications, poster presentations and abstracts at academic and professional conferences, discussion with peers, social media and the Global eHealth Unit website. The findings of this study will also inform JSM-B's PhD thesis.

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Contributors JSM-B developed the study protocol, and has been in charge of obtaining ethics and governance approvals. He also drafted this manuscript. AKG guided the study design, randomisation procedures and data analysis plan. He also performed the sample size calculations. CM, JO and JC guided the development of the study protocol, and have been supervising the work of JSM-B. All the authors reviewed and approved this manuscript.

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Competing interests None declared.

Ethics approval National Research Ethics Service Committee South East Coast.

Provenance and peer review Not commissioned; externally peer reviewed.

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