

BMJ Open Predicting perinatal health outcomes using smartphone-based digital phenotyping and machine learning in a prospective Swedish cohort (Mom2B): study protocol

Ayesha M Bilal ^{1,2}, Emma Fransson,^{3,4} Emma Bränn,³ Allison Eriksson,^{2,3} Mengyu Zhong,^{2,5} Karin Gidén,³ Ulf Elofsson,^{1,3} Cathrine Axfors,³ Alkistis Skalkidou,³ Fotios C Papadopoulos¹

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AS and FCP contributed equally.

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For numbered affiliations see end of article.

Correspondence to

Ayesha M Bilal;
ayesha.bilal@neuro.uu.se

ABSTRACT

Introduction Perinatal complications, such as perinatal depression and preterm birth, are major causes of morbidity and mortality for the mother and the child. Prediction of high risk can allow for early delivery of existing interventions for prevention. This ongoing study aims to use digital phenotyping data from the Mom2B smartphone application to develop models to predict women at high risk for mental and somatic complications.

Methods and analysis All Swedish-speaking women over 18 years, who are either pregnant or within 3 months postpartum are eligible to participate by downloading the Mom2B smartphone app. We aim to recruit at least 5000 participants with completed outcome measures. Throughout the pregnancy and within the first year postpartum, both active and passive data are collected via the app in an effort to establish a participant's digital phenotype. Active data collection consists of surveys related to participant background information, mental and physical health, lifestyle, and social circumstances, as well as voice recordings. Participants' general smartphone activity, geographical movement patterns, social media activity and cognitive patterns can be estimated through passive data collection from smartphone sensors and activity logs. The outcomes will be measured using surveys, such as the Edinburgh Postnatal Depression Scale, and through linkage to national registers, from where information on registered clinical diagnoses and received care, including prescribed medication, can be obtained. Advanced machine learning and deep learning techniques will be applied to these multimodal data in order to develop accurate algorithms for the prediction of perinatal depression and preterm birth. In this way, earlier intervention may be possible.

Ethics and dissemination Ethical approval has been obtained from the Swedish Ethical Review Authority (dnr: 2019/01170, with amendments), and the project fully fulfils the General Data Protection Regulation (GDPR) requirements. All participants provide consent to participate and can withdraw their participation at any time. Results from this project will be disseminated in

Strengths and limitations of this study

- The study collects large-scale, temporally sensitive data regarding the user's behaviours in the real world.
- End users' feedback collected allows for app updates and improvements.
- The passive data collection is expected to have lower attrition rate.
- The active data collection is prone to suffer from a higher attrition rate.
- There are high costs associated with recruiting participants and maintaining frontend and backend for the smartphone app.

international peer-reviewed journals and presented in relevant conferences.

INTRODUCTION

Optimal maternal health is important throughout pregnancy, childbirth and the postpartum period to ensure the full potential for the mother, infant and family to get a good start.¹ Two health conditions that are important to address in order to reach a goal of good maternal health are perinatal depression (PND) and preterm birth (PTB), both affecting about every 10th pregnancy worldwide.^{2,3}

Perinatal depression

PND is an episode of major depression with onset anytime during pregnancy and up to 4 weeks postpartum,⁴ although in research settings, a period of up to 1 year postpartum is often considered.⁵ Antenatal depression affects between 7% and 13% of pregnant women,⁶ and postpartum depression (PPD) is estimated to affect between 10% and 20% of all newly delivered mothers.²

while many women experience persistent depression throughout the perinatal period.⁷ PND is distinct from 'baby blues', which are commonly experienced symptoms of low mood and anxiety that subside within 2 weeks postpartum. PND is both emotionally and physically debilitating like major depression, with additional risks related to the pregnancy and birth, such as PTB, low birth weight, pre-eclampsia and placental abnormalities.^{8–10} Moreover, it is associated with retained maternal weight postpartum,¹¹ decreased breast feeding,^{12–14} poor maternal sleep¹⁵ and poor perinatal quality of life.¹⁶ PND can also compromise the critical mother–infant bond, as it affects the mother's caregiving abilities and adaptation to the maternal role,^{16 17} and has a long-term impact on the child's cognitive, emotional and behavioural development.^{18 19} Furthermore, PND can be characterised by the occurrence of self-harm thoughts, which are linked to long-term somatic and psychiatric morbidity,²⁰ and increased maternal mortality from suicide in the first year postpartum.²¹

The aetiology of PND is multifactorial, including biological, genetic, psychological and social factors, such as stressful life events, social support, domestic violence, childhood adversity, history of depression and anxiety, low self-esteem and even personality traits like resilience.^{15 22–24} Despite this knowledge, detecting PND remains a challenge for the healthcare system, with one review finding that around 30%–70% of cases go undetected and only 15% receive adequate treatment.^{25 26}

Current screening protocols include the Edinburgh Postnatal Depression Scale (EPDS)²⁷ during postpartum visits to assess risk of PND.²⁸ However, early detection of PND has remained challenging for many reasons, including inconsistencies in screening practices,²⁹ and failure to distinguish depressive symptoms due to their overlap with typical somatic experiences in the early postpartum period.³⁰ Furthermore, women may hesitate to seek care possibly because of the depression itself, but also stigma and fear of being judged as an imperfect mother, as well as concerns about antidepressant use during pregnancy and breast feeding.³¹ Besides, in-clinic screening frequently relies on retrospective self-reports of diagnostically relevant information, making it susceptible to errors and biases associated with autobiographical recollection.³²

Unquestionably, more efficient and effective methods for predicting PND in mothers are needed to enable early identification and intervention, thus improving prognosis, and reducing the burden of disease.³³ Previous studies that have attempted to develop predictive models of maternal depression primarily focus on the postpartum period only.^{34–39} Few have used social media fingerprints^{40–42} or biomarkers^{36 43} in their models, and these studies largely depended on psychometric self-reports and limited modalities. These drawbacks compromise the predictive power of the models and illustrate why multivariate, real-time and unobtrusive approaches to data collection and symptom monitoring must be encouraged to develop better predictive models.

Preterm birth

Among somatic pregnancy complications, PTB is a major cause of neonatal death, as well as of poor long-term health in children, affecting approximately 15 million babies worldwide each year.^{44 45} In Sweden, the PTB rate is about 6%,⁴⁶ which is a relatively low number compared with the international average of over 10%.⁴⁴ Like PND, the aetiology of PTB is multifactorial, including previous PTB, multifetal pregnancy, cervical insufficiency, intrauterine infections, vaginal bleeding in the second trimester, in-vitro fertilisation, primiparity, as well as maternal antenatal stress and depression.^{47–50} In fact, many risk factors overlap between PND and PTB, such as childhood traumatic events or maltreatment, stressful life events, being single or lacking social support, being overweight, smoking and low socioeconomic status.^{51 52} Inflammation has been suggested as a possible underlying pathway for both depression and preterm delivery.⁵³

There are evidence-based interventions for preventing or delaying PTB to optimise birth outcomes, such as smoking cessation, progesterone therapy, cerclage in women with cervical insufficiency or antibiotics.^{51 54} However, a major obstacle for the success of these interventions is the aetiological heterogeneity of PTB, which makes it extremely challenging to identify women at high risk. In fact, two-thirds of women who experience PTB do not present with any risk factors at all.⁵⁵ Available biological diagnostic tests for PTB (such as fetal fibronectin) lack sufficient positive prediction values.⁵⁶ Screening for cervical length is performed in Sweden for women with a history of PTB; however, this is not helpful in primiparous women.⁵⁷

It can be concluded that no single biomarker is sufficient for prediction; multimodal data, including psychosocial and behavioural factors, should, therefore, be the focus of prediction efforts.

Digital phenotyping and big data

Digital devices like smartphones allow us to capture moment-by-moment, objective data regarding the patient's experiences and functions in non-clinical settings. This process, known as *digital phenotyping*,⁵⁸ allows us to collect two kinds of data: *active data* and *passive data*. Active data refer to data that require user input, such as surveys and voice recordings. Passive data refer to automatically collected data from smartphone sensors and activity logs, which can be used to infer the user's mobility and sleep patterns, digital social activity, smartphone usage patterns, and even affective and cognitive changes.

The Mom2B smartphone app is developed using the Beiwe research platform (www.beiwe.org) from the Harvard School of Public Health. It can allow us to capture digital phenotyping data during the perinatal period with greater efficiency and temporal sensitivity as data collection occurs continuously and in real-world contexts, which minimises the risk of recall biases. Such apps could also be integrated into the mother's perinatal care plan. One drawback of smartphone-based data

collection, in general, is that attrition rate increases with longer follow-up times⁵⁹; however, this can at least in part be compensated for by the continuous collection of passive data.

In fact, one of the biggest advantages of smartphone-based digital phenotyping is the ability to collect multivariate, high-volume data, known commonly as big data.⁶⁰ Big data are excellent for healthcare research since it can facilitate a unique insight into risk factors and the development of better diagnostic frameworks⁶¹; however, the literature on big data approaches for psychiatric conditions, particularly perinatal mental health, is limited.⁶² Nordic countries are in the forefront in this respect—with the Danish National Birth Cohort,⁶³ the Norwegian Mother, Father and Child Cohort (MoBa)⁶⁴ and Autism Birth Cohort (ABC)⁶⁵ studies, and the Swedish Biology, Affect, Stress, Imaging and Cognition (BASIC) cohort study⁶⁶—due to the availability of nationwide registers with comprehensive personal and medical information for all pregnant women in these countries. Nonetheless, register data, while valuable, lack the multitude, continuity and veracity offered by digital phenotyping.⁶⁷ Furthermore, studies derived from these cohorts have largely relied on traditional statistical methods, which are limited in their ability to scale to large data sets and identify more subtle patterns in data.⁶⁸

To date, few studies have applied digital phenotyping for prediction of psychiatric conditions, such as relapse in schizophrenia⁶⁹ and severity of mood episodes in bipolar disorder.⁷⁰ In the context of PND, while smartphone apps are widely used, their application has been largely focused on screening and intervention.⁷¹ Only two studies have applied digital phenotyping for predicting PPD.^{34 72} While these studies have reported encouraging results, their predictive ability is compromised due to limited modalities (using only active data in the form of questionnaires), infrequent measurement points and usage of more traditional statistical methods. The Mom2B study combines nationwide health and pregnancy register data with active and passive data collected through smartphone-based digital phenotyping to objectively monitor indicators of PND in non-clinical contexts.

In order to harness the full potential of big data, more advanced analytical methods, such as machine learning (ML) and deep learning (DL), are ideal. ML is an artificial intelligence approach that refers to various methods of enabling an algorithm to identify and learn intricate patterns in data to predict outcomes.⁷³ Modern ML methods, such as deep neural networks (DNNs), are uniquely suited to analysing big data sets as they can detect complex, high-dimensional interactions and structured information, without guidance, that can then be used to train predictive algorithms. DNN models are comprised of multiple ‘hidden’ processing layers, inspired by biological neural networks, consisting of a series of interconnected nodes that resemble neurons.^{74 75}

Over the last decade, there has been a steady increase in the use of DL methods in medicine.⁷⁶ However, few

studies have used ML for diagnosis or risk assessment in psychiatry, and those that do are often limited by modest sample sizes and modalities, or from using only traditional ML techniques.^{73 77} To our knowledge, Mom2B is the first study to adopt a big data approach and use multimodal digital phenotyping with advanced ML techniques to develop predictive algorithms for PND and PTB.

Objectives

Using large-scale, multimodal data collected through the Mom2B smartphone app, together with health and pregnancy information from national registers, the primary aim of this study is to assess the accuracy of advanced ML and DL methods in predicting development of PND (1) in the third pregnancy trimester, using data from the first trimester, and (2) during the early and late postpartum period, using data collected throughout pregnancy and childbirth.

A secondary aim of this study is to apply advanced ML and DL techniques using the multimodal data set to predict the risk of PTB.

METHODS AND ANALYSIS

Cohort description

Mom2B (www.mom2b.se) is a national ongoing smartphone app-based study; the app was launched at the end of November 2019 to App Store and Google Play. All Swedish-speaking women above the age of 18 owning a smartphone, who are either pregnant or within 3 months postpartum, are eligible to participate by registering and providing consent in the Mom2B app. Participating women are also asked for optional consent to be contacted for additional research studies within and from outside the Mom2B project (see online supplemental appendix A). Participant data are then linked to psychiatric and somatic health-related and pregnancy-related information available from Swedish national registers.

We aim to recruit at least 5000 participants with completed outcome measures. Due to the complexity of ML methods, it is not possible to perform any traditional test of statistical power. However, based on previous studies^{59 78 79}, and conferring with experts in artificial intelligence, we estimated that this approximate number would give us enough material to build robust prediction models while accounting for attrition and the prevalence rates of the outcomes.

Information about the study is being disseminated on social media, and through posters and brochures sent to primary and maternal care centres across the country. Figure 1 illustrates Mom2B recruitment, data collection and opt-outs. Table 1 outlines participant characteristics of the existing Mom2B cohort based on users who have contributed relevant data, along with similar characteristics in the general Swedish population.

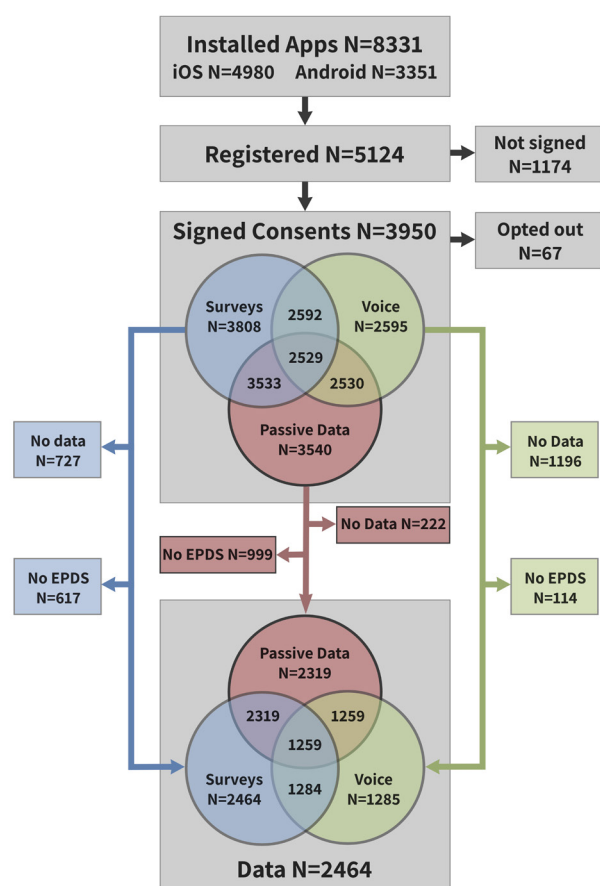


Figure 1 From top to bottom, the grey content blocks in the main column represent installed apps (downloads of the Mom2B app by unique users from either App Store (iOS) or Google Play (Android)), registered users (individuals who have submitted registration information in the app), signed consents (registered users who have consented to contributing data, and signed these consents electronically) and, finally, data (participants with signed consents who, at minimum, have completed the Edinburgh Postnatal Depression Scale (EPDS²⁷) at least once). The latter two blocks also illustrate the signed consents and available data, respectively, by type of data (survey, voice and passive data). The intersections of the Venn diagrams are non-exclusive, meaning that the number count in the intersection of surveys and passive data, for example, can include individuals who have also contributed to voice recordings. This flow chart reflects data last downloaded on 6 September 2021.

Data collection

The Mom2B app collects three types of data: survey data, audio recordings, as well as passive data. Data can be collected from the first week of pregnancy, and up until week 52 after birth. Only data that participants have consented for are collected from the time they register to the study, and they can change their consent preferences anytime in the app if they wish to stop.

Surveys and questionnaires

The Mom2B app delivers a range of both validated and self-developed questionnaires two to three times per week on average, with a mean of five questions per survey. These

questionnaires are used to collect information regarding the participant's mental and physical well-being, and history, personality, relationships, as well as perinatal and parenthood experiences. They include the EPDS, a 10-item self-report screening tool with good psychometric properties,^{80 81} used as the primary outcome measure in this study to assess depressive symptoms throughout the study period. A summary of the timeline of validated and self-developed instruments, along with the number of occurrences of the survey throughout the study period, can be found in figures 2 and 3, respectively.

Voice recordings

Voice acoustic qualities, such as pitch, speed, timing and timbre, have been used in previous research to successfully distinguish depressed from non-depressed individuals.^{82–85} To collect voice data, the Mom2B app sends out a voice recording task asking the participant to record reading simple texts, numerical sequences or vocalisations every 2–4 weeks.

Passive data collection

Passive data that the user has provided consent for are continuously collected via the Mom2B app throughout the study period, and are used to infer the user's behavioural patterns. Some of these features are collected differently for iOS and Android users. The feature modalities are briefly explained below.

Mobility

Correlations have been demonstrated between a patient's geographical movement patterns and changes in depressive symptoms.^{86 87} Sixty seconds of Global Positioning System (GPS) data are continuously collected every 10 min. Accelerator data are collected when the motion exceeds a certain threshold; motion activities, including being stationary, walking, running, cycling and movement in a vehicle, are recorded when the state changes. For iOS, we collect device motion to provide more detailed motion sensor data.

Data usage

Internet usage is the main feature of data usage. Various patterns of internet usage have been identified in relation to depression in different populations,^{88 89} but not among women with PND. The Mom2B app records the accumulated upload and download rates together with timestamps. Another feature, reachability, records timestamped smartphone connectivity—whether the phone is connected to cellular network, Wi-Fi or neither. It also records a Wi-Fi log for Android phone, which includes anonymised Media Access Control (MAC) addresses' frequencies and Received Signal Strength Indicator (RSSI) of available wireless networks in the area.

Smartphone usage

General smartphone use has been found to correlate with sleep quality and depression.^{90 91} Phone power state, combined with mobility parameters, can reflect individual

Table 1 Sociodemographic characteristics, pregnancy history and birth outcomes on participants in the Mom2B study and the general population of pregnant women in Sweden

Characteristics	Mom2B (2020–2022) (n=3909)*			Sweden (2019)†	
	Available data (n)	Missing data (n)	n (%) or mean±SD	Available data (n)	n (%) or mean
Maternal age (years)	3430	479	31.2±4.4	113 816	30.7
Country of origin	3441	468		112 530	
Sweden			3177 (92.3)		78 033 (69.3)
Nordic countries except Sweden			40 (1.2)		1280 (1.1)
Europe except Nordic countries			116 (3.4)		9172 (8.2)
Outside Europe			108 (3.1)		24 045 (21.4)
Education	3444	465		107 711	
≤12 years			744 (21.6)		48 793 (45.3)
Post-secondary education			2700 (78.4)		58 918 (54.7)
Employment before pregnancy	1677	2232		113 147	
Working/student/parental leave			1626 (97)		103 967 (91.9)
Unemployed/sick leave			51 (3)		9180 (8.1)
Smoking 3 months before pregnancy	3041	868	441 (14.5)	110 991	11 765 (10.6)
BMI before pregnancy (kg/m ²)	3353	556	25.5±5.3	108 929	
<18.5			70 (2.1)		2783 (2.5)
18.5–25			1815 (54.1)		59 384 (54.6)
25–<30			923 (27.5)		29 636 (27.2)
≥30			545 (16.3)		17 126 (15.7)
Primiparous	3268	641	1188 (36.4)	113 816	48 473 (42.5)
Caesarean section	1356	639‡	238 (17.5)	114 757	20 312 (17.7)
Preterm delivery (<week 37)	3311	598	190 (5.7)	116 071	6502 (5.6)

Percentages are given in relation to available data from women.

*Data downloaded on 1 February 2022.

†Data retrieved from the Swedish Medical Birth Register and Swedish National Board of Health and Welfare from 2019.

‡Calculated using the confirmed number of women in the postpartum period only.

BMI, body mass index.

behaviour like sleep patterns.^{92 93} To keep track of the use of the smartphone device, data are collected on screen activity, charging status and device reboot.

Social media activity

Social media behaviour has also been proven useful in detecting mental states. It has also been shown that reduced social activity on Facebook predicted symptoms of PPD.⁴¹ Collected data consist of simple behavioural measures, such as posting, commenting or liking, together with their timestamps. Notably, we only measure activity levels, not information related to the text or image

content of that activity, and participants are made aware of this when providing consent.

Survey metadata

App-based surveys make it possible to also collect metadata. This kind of behavioural metadata may contain clinically relevant information related to attention, processing speed and working memory capacity, and even any deterioration of psychiatric symptoms.^{58 94} We collect data on the time a survey was opened, time taken to answer each question and fully complete a survey, as well as any changes made in survey responses.

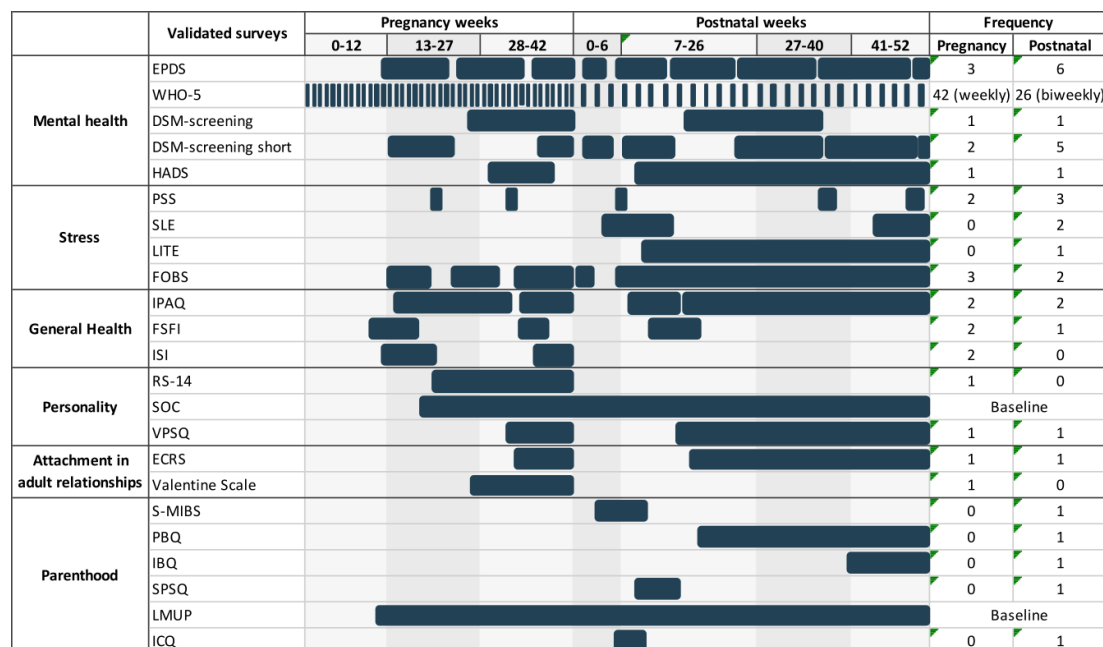


Figure 2 Timeline of validated instruments administered in the Mom2B study during pregnancy and postpartum periods, and the number of occurrences for each instrument throughout the study period. Surveys become available to users for varying periods of time and will disappear once completed or when their period of availability is over. EPDS, Edinburgh Postnatal Depression Scale²⁷; WHO-5, WHO-5 Well-Being Index¹⁰⁶; DSM-screening, Diagnostic and Statistical Manual of Mental Disorders, 5th Edition, criterion for depression; DSM-screening short is a shortened version of the DSM-screening with selected questions chosen by the research team; HADS, Hospital Anxiety and Depression Scale¹⁰⁷; PSS, Perceived Stress Scale¹⁰⁸; SLE, Stressful Life Events¹⁰⁹; LITE, Lifetime Influence of Traumatic Experiences¹¹⁰; FOBS, Fear of Birth Scale¹¹¹; IPAQ, International Physical Activity Questionnaire¹¹²; FSFI, Female Sexual Function Index¹¹³; ISI, Insomnia Severity Index¹¹⁴; RS-14, Resilience Scale¹¹⁵; SOC, Sense of Coherence¹¹⁶; VPSQ, Vulnerability Personality Style Questionnaire¹¹⁷; ECRS, Experience in Close Relationships Scale¹¹⁸; Valentine Scale (relationship with your partner)¹¹⁹; S-MIBS, Swedish Mother to Infant Bonding Scale¹²⁰; PBQ, Postpartum Bonding Questionnaire¹²¹; IBQ, Infant Behavior Questionnaire¹²²; SPSQ, Swedish Parenthood Stress Questionnaire¹²³; LMUP, London Measure of Unplanned Pregnancy¹²⁴; ICQ, Infant Characteristics Questionnaire.¹²⁵

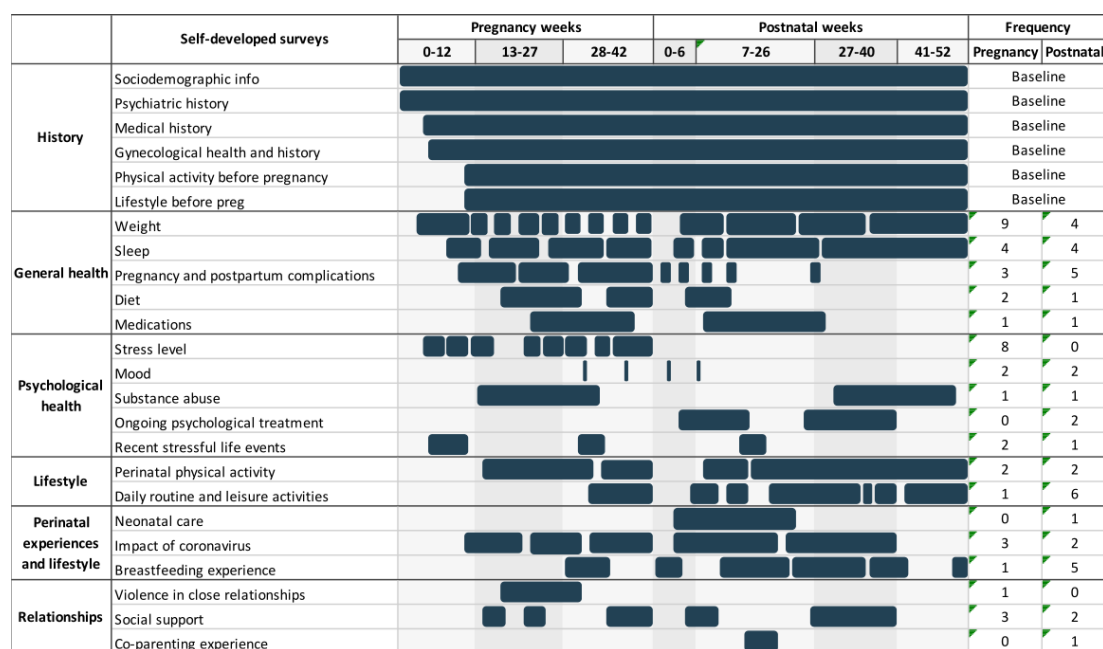


Figure 3 Timeline of self-developed surveys administered in the Mom2B study during pregnancy and postpartum periods, and the number of occurrences for each instrument throughout the study period. Surveys become available to users for varying periods of time and will disappear once completed or when their period of availability is over.

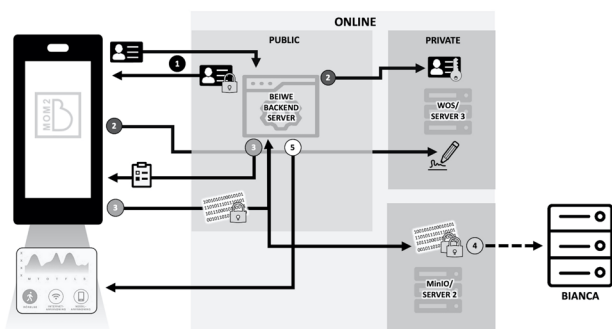


Figure 4 Flow of data from user to servers for storage and analysis. Data pass through secure servers accessible only by authorised members of the Mom2B team, and can be decrypted for analysis in Bianca when needed.

National register data

Supplementary information will be accessed via the following Swedish national health and quality registers: the Medical Birth Register, the Pregnancy Register, the National Patient Register, the Prescribed Drug Register and population censuses from Statistics Sweden. The accessed information includes records of perinatal complications such as PND, PTB or any other complications considered important risk factors for the study outcomes, such as gestational diabetes, gestational hypertension, pre-eclampsia, prolonged delivery, severe lacerations, postpartum haemorrhage, induction of delivery, instrumental vaginal delivery, caesarean section and small for gestational age.

Further, the mother's weight at enrolment in maternity care and at aftercare visits; calculated date of birth from last menstrual period and from ultrasound; and information on previous miscarriages, previous abortions, chronic diseases, fear of childbirth, involuntary infertility, gestational age at enrolment in maternity care and fetal diagnostics will be obtained from the Medical Birth Register. Retrieved information also includes variables regarding the background, health and lifestyle of the participant for validation purposes of our self-report questionnaires, as well as psychiatric and somatic morbidity for up to 15 years after childbirth.

Data flow and storage

Figure 4 illustrates the data flow and storage process as follows:

- ▶ Participants register to the study via the Mom2B app using their Swedish Social Security number, which is encrypted in the device using a private key provided by the Beiwe backend server, and replaced by a random, pseudoanonymised code number.
- ▶ The decryption key, together with the participant consent information and electronic signatures, is stored in a private, write-only server at Uppsala University.
- ▶ The app fetches surveys and voice recording tasks, and uploads data from participants to the backend server, where it is encrypted and sent to MinIO, a

secure, cloud-based storage, where another layer of encryption is added. Passive data collected from the phone follow the same path.

- ▶ From MinIO, all data are sent to Bianca, a private offline server, in both encrypted and decrypted forms for storage and analysis, respectively.
- ▶ The app provides weekly reports based on participant activity and fetches health-related information relating to the perinatal period the user is in, as well as frequently asked questions about the study and perinatal health.

Preliminary data analysis strategy

The Mom2B data set contains different modalities, including audio data, sensor data and survey data, which will be analysed separately and then combined. We plan to use both traditional ML and DL techniques in order to determine reliable predictors of PND and develop accurate predictive algorithms, and will report our findings following the best fit current guidelines, such as the Transparent Reporting of a Multivariable Prediction Model for Individual Prognosis or Diagnosis (TRIPOD) statement.⁹⁵

Feature engineering

To handle these multimodal data, we will extract features for these modalities separately using traditional feature engineering as well as DL techniques.^{96 97} An example of traditional feature engineering uses trajectories,⁸⁷ which use mobility features, such as number of significant places visited, maximum distance and SD. However, DL can be used for extracting features in many other modalities, such as social media and audio data, which are not investigated widely in the area.

Feature selection and model selection

To analyse the multimodal Mom2B data set, we will start with each modality separately. To reduce the possibility of potential overfitting, given the numerous features in our data set, we will use recursive feature elimination to obtain the optimal set of variables for further model development.

Previously, logistic regression, support vector machine, random forests, XGBoost and neural networks have been the most commonly used and efficient ML algorithms for prediction of PND.⁹⁸ An advantage of using such traditional ML methods is to give us a feature importance ranking, allowing us to identify stronger predictors. Using DL to analyse digital phenotyping data for evaluating risk of depression is a relatively novel approach compared with traditional ML models.⁷³ DL models have been shown to outperform traditional ML in various tasks involving complex data sets,^{99 100} and can be combined with traditional ML in multimodal data mining tasks to further improve performance.⁹⁷ We will test and select the best performing ML models for each modality and determine strong predictors of PND.

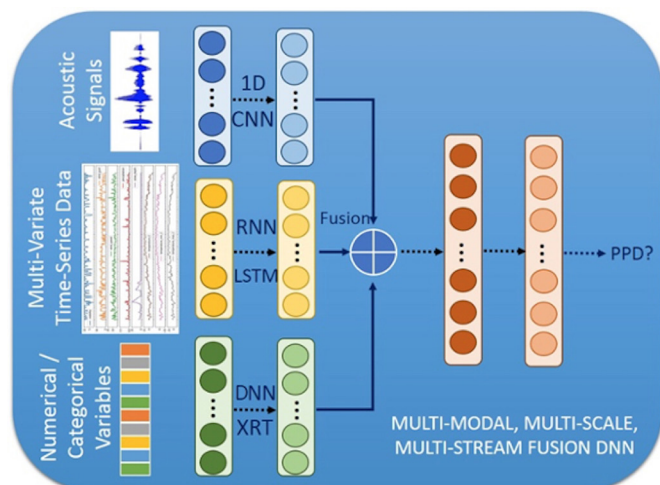


Figure 5 A multimodal machine learning model for peripartum depression (PPD) diagnosis. The extracted features can be classified into three categories: acoustic signals, time series features and categorical features. We can then determine the most suitable model for each category. For example, for acoustic signals, we would apply convolutional neural network (CNN); for time series data, we would apply recurrent neural network (RNN) such as long short-term memory (LSTM); and for numerical variables, we would apply deep neural networks (DNNs) such as transformers, or traditional models like extremely randomised trees (XRT), gradient boosted trees, etc. These models can yield high-dimension representations of multimodal features. After feature fusion, the integrated features will be fed into another neural network for prediction.

Multimodal computational model

The multimodal data we collect are in different scales, dimensions and formats, which need to be harmonised before prediction.¹⁰¹ Different models are better suited to perform on different modalities. To handle this complexity in multiple data modalities, we consider modality fusion during the development phase.

One example of a multimodal ML model is shown in figure 5. The model is designed to detect potential depressive episodes based on multimodal data collected in the Mom2B app. Preprocessed data in three modalities are fed into models and intermediate representations are then fused together and fed in as input features of a classification model.

Evaluation metrics

Our data will be split into a training data set, for analysis, and a test data set, to assess model performance. We consider using multiple evaluation metrics including area under the receiver operating characteristic curve, specificity, sensitivity, positive predictive value, negative predictive value, balanced accuracy and F1 score, as these measurements vary in importance according to the setting and goal of the final algorithm. Thereby, we can compare the performance of traditional ML with DL, and the different assemblies of models, from different perspectives in the context of prediction of PND.

Patient and public involvement

A qualitative study is planned for exploring the attitudes and concerns of participating women towards the Mom2B app. Furthermore, an online survey will be sent to women who have had no recent activity on the app or withdrawn participation from the study. Direct contact with end users and the ability to make changes to the app based on their feedback can enhance user experience and increase engagement. A representative from Mamma till Mamma, a non-profit organisation in Sweden focused on perinatal mental well-being, serves on our advisory board. The organisation has been involved in the piloting of our study and design of questionnaires, and currently supports us with recruitment. We plan to involve them in the dissemination of study results as well.

Substudies

In addition to predicting PND and PTB, the rich data collected from the Mom2B cohort will also be used to investigate further questions, mainly regarding the health of pregnant and postpartum women. Other planned areas of research are regarding the impact of early mother–infant separation and neonatal intensive care of the baby on the well-being of the mother; and sexual function and its potential correlates to depression and anxiety in the perinatal period.

Maternal depression and well-being during the COVID-19 pandemic

In the beginning of 2021, data collected from 1577 participants were used to assess depressive and anxiety symptoms, as well as well-being and life changes in pregnant women in Sweden during the COVID-19 pandemic (from February 2020 to March 2021).¹⁰² The Mom2B app enabled gathering psychiatric information at a national level during the pandemic, as well as passive data on mobility. Levels of perinatal affective symptoms and low well-being were elevated compared with previous years and to months with fewer cases. Similar apps can help healthcare providers and governmental bodies to monitor high-risk groups during crises in real time, as well as to adjust measures and the support offered.

ETHICS AND DISSEMINATION

Participants are informed about the aims of the study, and that the confidentiality and security of their data will be assured. All participants provide their consent to participate while registering to the study, and are informed that they can withdraw their participation at any time without giving a reason. Ethical approval has been obtained from the Swedish Ethical Review Authority (dnr: 2019/01170, with amendments) and the project fulfils General Data Protection Regulation (GDPR) requirements, including the processing, storage and protection of all data. Results will be continuously disseminated through international peer-reviewed journals, the project's website and social media channels, and presented in national and

international conferences. All publications will be open access.

DISCUSSION

Strengths and limitations

Besides the utility of digital phenotyping in combination with the advanced analytical methods planned to be used, other strengths of the Mom2B study include the involvement of participants. Statistics based on the WHO-5 Well-Being Index and behavioural data (movement, internet usage, sleep, etc) collected from participants are sent to the user, allowing them to follow their well-being and activity as an incentive for continued participation. Weekly informational reports regarding common experiences and concerns for both the mother and the child for that particular week of the perinatal period, based on information taken from 1177.se (Swedish health-care service), are available to users and allow them to easily stay informed. As per standard guidelines,¹⁰³ if participants receive a high score on the EPDS, they are prompted to contact their healthcare provider or emergency support services for support, and if unsure, they can contact the research team, which will help them find appropriate support for their needs. Continuous contact is maintained with participants until they find support.

The involvement of user organisations and an international advisory board further strengthens the study by increasing the feasibility, the use of state-of-the-art methods and the potential for high acceptance by the end users, which is especially important for future integration in regular clinical practice.

However, there are some limitations to acknowledge. Weekly reports and statistics are important in supporting and incentivising users, but it is possible they may influence users' responses to certain questionnaires. To account for this, we consider including how often they are checked by users as a feature within our models. Furthermore, our app is available only in Swedish, which excludes a number of otherwise eligible participants, and the high costs for maintaining the technical infrastructure in the frontend and backend of the app require considerable funding. Attrition is also an issue, especially with data that require active input from users. While we can attempt to combat this by improving the app based on user feedback, it is important to consider that attrition might also reflect the worsening of symptoms and be a predictor per se of clinical deterioration. It will be important to distinguish such participants and determine how to use attrition as a predictor variable.

Future perspectives

We are at the beginning of the smartphone-based research era, and future possibilities seem numerous. We intend to develop the Mom2B app in other languages, including English, to expand to a more diverse and wider population. If the app succeeds in developing good predictive models for PND, the research team anticipates that the

app could be further developed to include evidence-based interventions.¹⁰⁴ Furthermore, since PND is much less understood in co-parents and improving the other parent's mental well-being is conducive to the health of the mother and the children as well,¹⁰⁵ the app could be further developed to study co-parental PND. The Mom2B research team plans to further adapt the app to other research topics such as teenage and student mental health, and prediction of new episodes or self-harm in major depression.

Author affiliations

¹Department of Medical Sciences, Uppsala University, Uppsala, Sweden

²Centre for Women's Mental Health during the Reproductive Lifespan (Womher), Uppsala University, Uppsala, Sweden

³Department of Women's and Children's Health, Uppsala University, Uppsala, Sweden

⁴Centre for Translational Microbiome Research, Department of Microbiology, Tumor and Cell Biology, Karolinska Institute, Stockholm, Sweden

⁵Department of Information Technology, Uppsala University, Uppsala, Sweden

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ORCID iD

Ayesha M Bilal <http://orcid.org/0000-0002-7349-8765>

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Translated English version of participant information and consent form given digitally to participants via the Mom2B app. See further below for original Swedish version.

PARTICIPANT INFORMATION MOM2B

We want to ask you if you want to participate in a research project. Now you will receive information about the project and what it means to participate. There is a lot of text to read, but the text can also be read again in the app.

At the end of the information, you will have to give consent to the study. You can go into settings at any time and change your consents.

If you have questions about the study, you are welcome to contact us by e-mail:

mom2b@kbh.uu.se.

WHAT IS THE PROJECT AND WHY DO YOU WANT ME TO PARTICIPATE?

Postpartum depression is when women experience depression during pregnancy and up to one year after the baby is born. About 12% of all mothers experience postpartum depression. This can lead to suffering and problems for the affected mother and her children. There are effective treatment options. If research finds better ways to identify women at high risk for postpartum depression, more people could receive care early.

During pregnancy and childbirth, the mother and baby can have complications. An example is premature birth, when the baby is born before 37 weeks of pregnancy. Children born prematurely are at greater risk of becoming ill as newborns. In Sweden, about 5% of all births are premature. If the research provides better knowledge to predict which women are at higher risk of being affected, some complications could perhaps be prevented. With the help of smartphones, it is possible to collect information about people's everyday lives. Examples of measurements are how much a person moves and how active they are on social media. Such measurements have in previous studies been able to predict mental state, but it has not been studied in pregnant women or new mothers. No one has studied whether the measurements can predict complications during pregnancy and childbirth.

THE PURPOSE OF THIS STUDY

The purpose is to use data collected through this app, MOM2B, to improve the detection of women who are at higher risk of developing mental or physical illness during pregnancy and childbirth.

HOW DOES THE STUDY WORK?

You can participate in the study if these three points apply to you:

- I am pregnant or have given birth less than three months ago.
- I am currently using my own smartphone.
- I am at least 18 years old.

At the end of this information section, you can choose which parts of the study you wish to participate in. As a participant, you can choose to contribute:

- To answer questions via this app about your health, lifestyle and background as well as about the child. The questions are compiled by the research team and examine possible risk factors. Questions are sent out approximately 1-2 times per week and usually contain about 5 questions, a maximum of about 15 questions.

- Information about your general mobile use. This means how much internet you use, how much of the mobile processor you use, how often you have access to the internet and how often the charger is connected. The information is collected passively without you having to enter it.
- Information on how much you move. Then the phone's GPS and accelerometer are used. We do not save any specific GPS coordinates, but only information about how you moved in relation to a randomly selected location (eg 100m north, then 50m east, etc.). The information is collected passively without you having to enter it.
- Information about how active you are on social media. What is measured is the number of times someone "likes" something you posted, the number of times you make a status update and the number of times you send a message. Only the number of activities is saved, not the content. What you like or what text you wrote to whom is not saved. The information is collected passively without you having to enter it. The social media that is included right now is Facebook. There may be more (such as Instagram, LinkedIn, Twitter and Snapchat) but then you will get a new question about participation for these.
- To record the sound of your voice when reading texts or numbers. With the help of these audio recordings, we examine e.g. speech tempo and voice mode.
- Information from national health registers and care quality registers about your health, lifestyle and background from one year before the birth to 15 years after the birth. Examples of information that can be obtained are diagnoses you receive in contact with healthcare, level of education, income and employment. The information is collected via the National Board of Health and Welfare and Statistics Sweden.

POSSIBLE CONSEQUENCES AND RISKS OF PARTICIPATING IN THIS STUDY

Answering questions about mental health can involve a risk of emotional strain as some questions may be perceived as sensitive. You choose which questions you want to answer and if you need support, you can contact us for guidance.

If your questionnaire responses indicate severe symptoms of depression, you will receive an automatic message with information on how to seek care. Because your answers are coded and encrypted, we cannot contact you through the app, but you will have the opportunity to enter your phone number so that we can call you during office hours.

Although the research group takes measures to achieve very high data security (such as storing data under code instead of identity, encrypting data directly when collecting and storing data on Uppsala University's servers with a high level of physical and electronic security), it is impossible to completely rule out data intrusion. If data breach is suspected, it will be handled in accordance with current legislation.

ARE THERE ANY BENEFITS?

As a participant, you get access to a knowledge bank in the MOM2B app about health during pregnancy and after childbirth. You also get the opportunity to submit proposals for new topics to the knowledge bank. In addition, participation will not provide any immediate benefits.

WHAT HAPPENS TO MY TASKS?

The project will collect and register information about you. Here we describe how we protect your information.

Code number

You will automatically receive a code number so that no information is stored together with your identity. When we analyze the data and report the results, you will remain anonymous. Responses to questionnaires will be presented so that individuals' responses cannot be traced. The link between your code number and your identity is stored in an isolated location, separate from the information collected.

Encryption

Immediately when data is collected, an encryption is performed. This means that the information is made illegible. Only with the encryption key will the information be readable. All transport of data over networks is done with an additional encryption. The encryption key is stored on a storage unit without a network connection, which is stored locked in a safe at Uppsala University.

Storage of data

The collected information is stored coded and encrypted on a server at Uppsala University indefinitely. Prior to our analyses, the encryption will be unlocked, but the information will remain coded. Only researchers in our research group will have access to the data. Our main analysis, so-called machine learning, is done on computers in a locked space at Uppsala University that has no network connection. All in all, this means that the risk of data intrusion is greatly minimized.

You can choose whether some of your survey answers are to be presented in simplified form on your smartphone under "Statistics". No other data is stored on the phone itself. Identifiable personal data will never be shared with any third party other than the administrators at the National Board of Health and Welfare, Statistics Sweden and the quality of care register, who retrieve information from health registers for the participants who have consented to this. Immediately after the link, all data is deidentified again by coding.

Your answers and your results will be processed so that unauthorized persons cannot take part in them. Uppsala University is responsible for your personal data. According to the EU Data Protection Regulation, you have the right to access the information about you that is handled in the study free of charge and, if necessary, have any errors corrected. You can also request that information about you be deleted and that the processing of your personal data be restricted. If you want to take part in the information, you should contact the principal researcher, Alkistis Skalkidou, telephone: 070 167 97 78 or e-mail: mom2b@kbh.uu.se. The Data Protection Officer can be reached by e-mail: dataskyddsbud@uu.se. If you are dissatisfied with how your personal data is processed, you have the right to report this to the Swedish Data Inspectorate, which is the supervisory authority. The answers to surveys will be compiled statistically in anonymous form and presented so that individuals' answers cannot be traced.

How do I get information about the results of the study?

Results from the project will be presented on our website <https://www.mom2b.se/>, in scientific articles and at national and international conferences. The results of the study will present the participants' answers in groups and your individual answers will not be traceable to you.

Insurance and compensation

As a participant in this study, you are covered by the patient injury association. No compensation will be given for participating in the study.

PARTICIPATION IS VOLUNTARY

Your participation is voluntary and you can choose to cancel your participation at any time.

If you choose not to participate or want to cancel your participation, you do not have to state why, nor will it affect your future care or treatment.

If you would like to cancel your participation, please contact the lead researcher, Alkistis Skalkidou (see below). You are also welcome to contact us at mom2b@kbh.uu.se if you have questions about the study.

Responsible for the study are:

Professor Alkistis Skalkidou

Department of Women's and Children's Health

Akademiska Sjukhuset, 751 85 Uppsala

Email: mom2b@kbh.uu.se

Phone: 070 167 97 78

STEP 1: CONSENT

I have received written information about the study and have had the opportunity to ask questions. I have access to the written participant information electronically in the MOM2B smartphone application at any time.

In the paragraphs that come now, you decide which parts of the study you want to participate in by clicking in the boxes for Yes or No.

I approve of my participation in the study. (You must answer 'Yes' to be able to participate in the study.)

☐ Yes ☐ No.

STEP 2: SURVEYS

I agree to answer questions about my health, lifestyle and background as well as the child's health.

☐ Yes ☐ No.

STEP 3: MOBILE USE

I agree that data about my general mobile use is stored. This means internet consumption (kb/s), processor usage (CPU%/s), internet access and whether my phone is charging or not.

☐ Yes ☐ No.

STEP 4: MOBILITY PATTERN

I agree that data is stored about how much I move. We want to record how much and how fast you move, but not exactly where. Only location data from an unknown point is saved, not your exact location. If you select "no", no location data, relative or exact, is saved.

☐ Yes ☐ No.

STEP 5: SOCIAL MEDIA

I agree that information about how active I am on social media is stored. This means the number of times I perform an activity (that someone likes one of your status updates or that you post a status update) but not the text or image content of the activity. Therefore, this does NOT mean information about who is contacted, what is written or what is liked.

☐ Yes ☐ No.

STEP 6: AUDIO RECORDING

I agree to record the sound of my voice when I read out predetermined texts and numbers.

☐ Yes ☐ No.

STEP 7: HEALTH REGISTERS

I agree that information is obtained from national health registers and care quality registers about my health, lifestyle and background from one year before to 15 years after the birth. Examples are diagnoses I get in contact with healthcare, level of education, income and employment.

☐ Yes ☐ No.

STEP 8: PARTICIPATION IN STUDIES

To improve the app and increase the possibilities for predicting postpartum depression and facilitate other studies in women's health, we would like to ask you if we can contact you at a later opportunity with a request for participation in other studies.

At that time, you always have the opportunity to decline the offer of participation.

Participation in other studies is completely voluntary and if you choose not to be contacted, you will not be negatively affected in any way. Select one or both of the following options to be contacted:

☐ I wish to be contacted for inquiries about participation in sub-studies within Mom2B

☐ I wish to be contacted for inquiries about participation in other relevant studies on women's health

SIGNATURE

To be able to use the app, we need your signature

First and last name

[Click here to sign](#)

Original Swedish version of participant information and consent form given digitally to participants via the Mom2B app.

DELTAGARINFORMATION MOM2B

Vi vill fråga dig om du vill delta i ett forskningsprojekt. Nu kommer du att få information om projektet och vad det innebär att delta. Det är mycket text att läsa men texten finns också att läsa igen i appen.

I slutet av informationen kommer du få lämna samtycke till studien. Du kan när som helst gå in i inställningar och ändra dina samtycken.

Om du har frågor om studien är du välkommen att kontakta oss på e post:

mom2b@kbh.uu.se.

VAD ÄR DET FÖR PROJEKT OCH VARFÖR VILL NI ATT JAG SKA DELTA?

Förlossningsdepression kallas det när kvinnor drabbas av nedstämdhet under graviditeten och upp till ett år efter att barnet fötts. Ungefär 12% av alla mammor får förlossningsdepression. Detta kan leda till lidande och problem för den drabbade mamman och hennes barn. Det finns effektiva behandlingsalternativ. Om forskningen hittar bättre sätt att identifiera kvinnor med hög risk för förlossningsdepression skulle fler kunna få vård tidigt.

Under graviditeten och vid förlossningen kan mamman och barnet råka ut för komplikationer. Ett exempel är för tidig födsel, när barnet föds innan 37 graviditetsveckor. Barn som föds för tidigt har större risk för att bli sjuka som nyfödda. I Sverige är cirka 5% av alla födselar för tidiga. Om forskningen ger bättre kunskap för att förutsäga vilka kvinnor som har högre risk att drabbas skulle en del komplikationer kanske kunna förhindras.

Med hjälp av smartphones går det att samla in information om människors vardagsliv.

Exempel på mätningar är hur mycket en person förflyttar sig och hur aktiv den är på sociala medier. Sådana mätningar har i tidigare studier kunnat förutsäga psykiskt mående, men det har inte studerats hos gravida kvinnor eller nyblivna mammor. Ingen har heller studerat om mätningarna kan förutsäga komplikationer under graviditeten och vid förlossningen.

SYFTET MED STUDIEN

Syftet är att använda data som samlas via den här appen, MOM2B, för att förbättra upptäckandet av kvinnor som har en högre risk att drabbas av psykisk eller fysisk sjuklighet under graviditet och förlossning.

HUR GÅR STUDIEN TILL?

Du kan delta i studien om dessa tre punkter stämmer in på dig:

- Jag är gravid eller har fött barn för mindre än tre månader sedan.
- Jag använder just nu min egen smartphone.
- Jag är minst 18 år gammal.

I slutet av det här informationsavsnittet kan du välja vilka delar av studien du önskar delta i. Som deltagare kan du välja att bidra med:

- Att svara på frågor via den här appen om din hälsa, livsstil och bakgrund samt om barnet. Frågorna är sammanställda av forskarteamet och undersöker möjliga riskfaktorer. Frågor skickas ut ungefär 1-2 gånger per vecka och innehåller som regel ca 5 frågor, max ca 15 frågor.

- Uppgifter om din allmänna mobilanvändning. Med detta menas hur mycket internet du använder, hur mycket av mobilens processor du använder, hur ofta du har tillgång till internet och hur ofta laddaren är inkopplad. Informationen samlas passivt utan att du behöver skriva in den.
- Uppgifter om hur mycket du förflyttar dig. Då används telefonens GPS och accelerometer. Vi sparar inte några specifika GPS-koordinater, utan bara information om hur du rört dig i förhållande till en slumpmässigt utvald plats (t.ex. 100m norrut, sen 50m österut, osv.). Informationen samlas passivt utan att du behöver skriva in den.
- Uppgifter om hur aktiv du är på sociala medier. Det som mäts är antalet gånger någon "gillar" något du postat, antalet gånger du gör en statusuppdatering och antalet gånger du skickar ett meddelande. Bara antalet aktiviteter sparas, inte innehållet. Vad du gillar eller vilken text du skrivit till vem sparas inte. Informationen samlas passivt utan att du behöver skriva in den. De sociala medier som ingår just nu är Facebook. Det kan komma att bli fler (så som Instagram, LinkedIn, Twitter och Snapchat) men då kommer du att få en ny fråga om deltagande för dessa.
- Att spela in ljudet av din röst när du läser upp texter eller siffror. Med hjälp av dessa ljudinspelningar undersöker vi t.ex. taltempo och röstläge.
- Uppgifter från nationella hälsoregister och vårdkvalitetsregister om din hälsa, livsstil och bakgrund från ett år före förlossningen till 15 år efter förlossningen. Exempel på information som kan inhämtas är diagnoser du får i kontakt med vården, utbildningsnivå, inkomst och sysselsättning. Uppgifterna hämtas via Socialstyrelsen och Statistiska Centralbyrån.

MÖJLIGA FÖLJDER OCH RISKER MED ATT DELTA I STUDIEN

Att besvara frågor om psykisk hälsa kan innebära en risk för känslomässig påfrestning eftersom vissa frågor kan uppfattas som känsliga. Du väljer själv vilka frågor du vill svara på och ifall du behöver stöd kan du kontakta oss för vägledning.

Om dina enkätsvar skulle tyda på allvarliga symptom på nedstämdhet kommer du att få ett automatiskt meddelande med information om hur du kan söka vård. Eftersom dina svar är kodade och krypterade kan vi inte kontakta dig genom appen, men du kommer att ha möjlighet att ange ditt telefonnummer så att vi kan ringa upp dig under kontorstid.

Trots att forskargruppen vidtar åtgärder för att uppnå mycket hög datasäkerhet (som att lagra data under kod istället för identitet, kryptera data direkt vid insamling och förvara data på Uppsala Universitets servrar med hög nivå av fysisk och elektronisk säkerhet) är det omöjligt att helt utesluta dataintrång. Om dataintrång mot förmodan skulle ske kommer det att hanteras enligt gällande lagstiftning.

FINNS DET NÅGRA FÖRDELAR?

Du som deltagare får tillgång till en kunskapsbank i MOM2B-appen om hälsa under graviditet och efter förlossning. Du får också möjlighet att skicka in förslag på nya ämnen till kunskapsbanken. Utöver detta kommer deltagandet inte ge några omedelbara fördelar.

VAD HÄNDER MED MINA UPPGIFTER?

Projektet kommer att samla in och registrera information om dig. Här beskriver vi hur vi skyddar dina uppgifter.

Kodnummer

Du får automatiskt ett kodnummer så att inga uppgifter lagras tillsammans med din identitet. När vi analyserar uppgifterna och redovisar resultaten kommer du att förbli anonym. Svar på enkäter kommer att presenteras så att enskilda personers svar inte kan spåras. Kopplingen mellan ditt kodnummer och din identitet lagras på en isolerad plats, separat från de insamlade uppgifterna.

Kryptering

Direkt när uppgifter samlas in görs en kryptering. Detta betyder att informationen görs oläslig. Bara med krypteringsnyckeln blir uppgifterna läsliga. Alla transporter av data över nätverk görs med ytterligare en kryptering. Krypteringsnyckeln sparas på en lagringsenhet utan nätverkskoppling, som förvaras inlåst i ett kassaskåp på Uppsala Universitet.

Förvaring av datauppgifter

De insamlade uppgifterna sparas kodade och krypterade på en server vid Uppsala Universitet på obestämd tid. Inför våra analyser kommer krypteringen att låsas upp, men uppgifterna förblir kodade. Bara forskare i vår forskargrupp kommer att ha tillgång till uppgifterna. Vår huvudsakliga analys, s.k. maskininlärning, görs på datorer i ett låst utrymme på Uppsala Universitet som inte har någon nätverksuppkoppling. Detta innebär sammantaget att risken för dataintrång kraftigt minimeras.

Du kan välja ifall vissa av dina enkätsvar ska presenteras i förenklad form på din smartphone under "Statistik". Inga andra uppgifter lagras på själva telefonen.

Identifierbara personuppgifter kommer aldrig att delas med någon tredje part förutom de handläggare på Socialstyrelsen, Statistiska Centralbyrån och vårdkvalitetsregister som hämtar information från hälsoregister för de deltagare som samtyckt till detta. Omedelbart efter sammanlänkningen avidentifieras all data igen genom kodning.

Dina svar och dina resultat kommer att behandlas så att inte obehöriga kan ta del av dem. Ansvar för dina personuppgifter har Uppsala Universitet. Enligt EU:s dataskyddsförordning har du rätt att kostnadsfritt få ta del av de uppgifter om dig som hanteras i studien och vid behov få eventuella fel rättade. Du kan också begära att uppgifter om dig raderas samt att behandlingen av dina personuppgifter begränsas. Om du vill ta del av uppgifterna ska du kontakta huvudansvarig forskare, Alkistis Skalkidou, telefon: 070 167 97 78 eller e-post: mom2b@kbh.uu.se. Dataskyddsombud nås på e-post: dataskyddsombud@uu.se. Om du är missnöjd med hur dina personuppgifter behandlas har du rätt att anmäla detta till Datainspektionen, som är tillsynsmyndighet. Svaren på enkäter kommer att sammanställas statistiskt i anonym form och presenteras så att enskilda personers svar inte kan spåras.

Hur får jag information om resultatet av studien?

Resultat från projektet kommer att presenteras på vår hemsida <https://www.mom2b.se>, i vetenskapliga artiklar och på nationella och internationella konferenser. Studiens resultat

kommer att presentera deltagarnas svar i grupper och dina individuella svar kommer inte att kunna spåras till dig.

Försäkring och ersättning

Som deltagare i denna studie täcks du av patientskadeföreningen. Ingen ersättning kommer att ges för att du deltar i studien.

DELTAGANDET ÄR FRIVILLIGT

Ditt deltagande är frivilligt och du kan när som helst välja att avbryta deltagandet. Om du väljer att inte delta eller vill avbryta ditt deltagande behöver du inte uppge varför, och det kommer inte heller att påverka din framtida vård eller behandling.

Om du vill avbryta ditt deltagande ska du kontakta huvudansvarig forskare, Alkistis Skalkidou (se nedan). Du är också välkommen att kontakta oss på mom2b@kbh.uu.se om du har frågor kring studien.

Ansvarig för studien är:

Professor Alkistis Skalkidou

Institutionen för Kvinnors och Barns hälsa

Akademiska Sjukhuset, 751 85 Uppsala

Epost: mom2b@kbh.uu.se

Telefon: 070 167 97 78

STEG 1: SAMTYCKE

Jag har fått skriftlig information om studien och har haft möjlighet att ställa frågor. Jag har när som helst tillgång till den skriftliga deltagarinformationen elektroniskt i smartphone-applikationen MOM2B.

I de stycken som kommer nu, bestämmer du vilka delar av studien du vill vara med på genom att klicka i rutorna för Ja eller Nej.

Jag godkänner mitt deltagande i studien. (Du måste svara Ja för att kunna vara med i studien.)

☐ Ja ☐ Nej

STEG 2: ENKÄTER

Jag samtycker till att svara på frågor om min hälsa, livsstil och bakgrund samt barnets hälsa.

☐ Ja ☐ Nej

STEG 3: MOBILANVÄNDNING

Jag samtycker till att uppgifter lagras om min allmänna mobil användning. Med detta menas internetförbrukning (kb/s), processoranvändning (CPU%/s), tillgång till internet samt om min telefon är på laddning eller inte.

☐ Ja ☐ Nej

STEG 4: RÖRELSEMÖNSTER

Jag samtycker till att uppgifter lagras om hur mycket jag förflyttar mig. Vi vill registrera hur mycket och hur snabbt du rör dig, men inte exakt var. Bara platsdata från okänd punkt sparas, inte din exakta position. Väljer du "nej" sparas inga platsdata, relativa eller exakta.

☐ Ja ☐ Nej

STEG 5: SOCIALA MEDIER

Jag samtycker till att uppgifter lagras om hur aktiv jag är på sociala medier. Med detta menas antal gånger jag utför en aktivitet (att någon gillar en av dina statusuppdateringar eller att du postar en statusuppdatering) men inte aktivitetens text- eller bildinnehåll. Detta innebär alltså INTE information om vem som kontaktas, vad som skrivs eller vad som gillas.

☐ Ja ☐ Nej

STEG 6: LJUDINSPELNING

Jag samtycker till att spela in ljudet av min röst när jag läser upp förutbestämda texter och siffror.

☐ Ja ☐ Nej

STEG 7: HÄLSOREGISTER

Jag samtycker till att uppgifter hämtas från nationella hälsoregister och vårdkvalitetsregister om min hälsa, livsstil och bakgrund från ett år före till 15 år efter förlossningen. Exempel är diagnoser jag får i kontakt med vården, utbildningsnivå, inkomst och sysselsättning.

☐ Ja ☐ Nej

STEG 8: DELTAGANDE I STUDIER

För att förbättra appen och öka möjligheterna till prediktion av förlossningsdepression samt underlätta för andra studier inom kvinnohälsa, önskar vi fråga dig om vi får kontakta dig vid ett eventuellt senare tillfälle med förfrågan om deltagande i andra studier.

Vid den tidpunkten har du alltid möjlighet att avböja erbjudandet om deltagande.

Deltagande i övriga studier är helt frivilligt och väljer du att inte bli kontaktad kommer du inte att påverkas negativt på något sätt. Markera ett eller båda av följande alternativ om du vill bli kontaktad:

☐ Jag önskar bli kontaktad för förfrågan om deltagande i sub-studier inom Mom2B

☐ Jag önskar bli kontaktad för förfrågan om deltagande i andra. relevanta studier om kvinnohälsa

UNDERSKRIFT

För att kunna använda appen behöver vi din underskrift

För- och efternamn

Tryck här för att skriva under