Abstract

Background: Early warning systems (EWS) have been widely adopted for use in maternity settings internationally. The idea in using these systems is early recognition of potential or actual clinical deterioration in pregnant or postpartum women, and escalation of care. Barriers to successful implementation and use of EWS, however, have been identified. If EWS are to be applied consistently, a greater understanding of the views and experiences of EWS from the perspectives of those using and applying EWS in maternity practice is needed. This protocol describes a qualitative evidence synthesis of maternity care providers’ (midwives, obstetricians, and allied maternity care professionals) views and experiences of EWS use and application in practice.

Methods: Studies will be included in the review if they report on maternity care providers use and application of EWS in any birth setting. Qualitative studies and studies of mixed methods design, where qualitative data can be extracted separately, will be included. To source relevant literature the electronic databases of MEDLINE, CINHAL, Web of Science Core Collection (incorporating Social Science Citation Index) and Maternity and Infant Care (MIDIRS), from date of inception, will be searched. The methodological quality of the included studies will be appraised using the 12-criteria of the assessment tool developed by the Evidence for Policy and Practice Information and Co-ordinating Centre. Thematic synthesis will be used for synthesising the qualitative data from included studies. The confidence in the findings will be assessed using the Grading of Recommendations Assessment, Development and Evaluation-Confidence in the Evidence from Reviews of Qualitative research.

Conclusions: The findings of this qualitative evidence synthesis may provide valuable information on the barriers, challenges, and facilitators for EWS use based on the experiences of those directly involved in EWS application in maternity care provision.
PROSPERO registration: CRD42021235137 (08/04/2021)

Keywords
Early warning systems, maternity early warning score, MEWS, clinical deterioration, systematic review, qualitative evidence synthesis.

This article is included in the Maternal and Child Health collection.

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Author roles: Smith V: Conceptualization, Methodology, Writing – Original Draft Preparation, Writing – Review & Editing; Cithambaram K: Methodology, Writing – Review & Editing; O'Malley D: Methodology, Writing – Review & Editing

Competing interests: No competing interests were disclosed.

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Introduction

Early warning systems (EWS) have been introduced in clinical care as a means of formalising the measurement of physiological variables (temperature, pulse rate, blood pressure, respiratory rate, alert status, etc.). Measurements are recorded and collectively assessed, generally using either a points-based scoring system (0, 1, 2, 3, etc.) or a colour-coded traffic light system (red, orange, green) based on each parameter measuring within or outside of agreed physiological norms. The idea in using these systems is early recognition of potential or actual clinical deterioration in pregnant or postpartum women, and escalation of care. Reports spanning more than a decade from professional bodies and other national maternity clinical guidelines recommend using EWS to identify the potential for clinical deterioration in women who are pregnant or postpartum. These systems have now been widely adopted for use in maternity settings internationally.

Although limited, some evidence exists which suggests that use of EWS may be useful in predicting maternal mortality, although less so for mortality. To affect accurate prediction, however, compliance in using EWS consistently in practice and according to recommended schedules, which may vary depending on location and clinical scenario (e.g. 1-hourly during labour, 4-hourly during routine postnatal care, etc.) is required. Studies have demonstrated significant variance in compliance rates with EWS use, ranging from below 50% to 100% across studies. Compliance in use also appears to diminish as a woman’s length of stay in hospital extends. For example, Allman noted that non-recording of observations on EWS ranged from 64% at two hours to 2% at seven hours. Helme also reported an increased rate of ‘poor’ recordings on EWS, from 11% at one-hour post-surgery to 27% at two hours and 91% between three and 24 hours postoperatively. Of further concern, 40% of maternal EWS scores in one audit were reportedly inaccurate.

Barriers to successful implementation and use of EWS have been identified. These include overlap with other charts, staff shortages, lack of training in EWS use, lack of support for EWS, delegating observation recording to maternity care assistants, too time consuming and a lack of validation. If EWS is to be applied consistently in maternity care for the purpose of early detection and management of clinical deterioration there needs to be enhanced understandings of the views and experiences of EWS from the perspectives of those applying EWS in practice. For this reason, we plan to perform a synthesis of qualitative data of maternity care providers’ views and experiences of EWS.

Aim

This paper describes a protocol for a qualitative evidence synthesis of maternity care providers’ (midwives, obstetricians, and allied maternity care professionals) views and experiences of EWS use and application in practice. The proposed review is registered with PROSPERO (CRD42021235137, 8th April 2021) and adheres to the Preferred Reporting Items for Systematic review and Meta-Analysis Protocols (PRISMA-P) reporting guidelines.

Methods

Inclusion criteria

We use the SPIDER (sample, phenomenon of interest, design, evaluation, and research type) acronym to define the criteria for including studies in the review as follows:

- Sample: Maternity care providers (MCPs) involved in caring for pregnant/postpartum women, of low and high risk, and in any birth setting. MCPs are defined as professionally qualified MCP, for example, midwife nurse, doctor, obstetrician and other allied MCPs.
- Phenomenon of Interest: EWS application/use in clinical practice
- Design: Qualitative studies of any design. Mixed methods studies, where the qualitative data are accessible separately, will also be considered for inclusion. Survey designs with open-ended response options that collect qualitative data may be considered for inclusion. The depth of qualitative data must be sufficient however, for surveys to be included; that is, surveys that report qualitative data in the form of exemplar quotes to support quantitative ‘counts’ or where the data are briefly summarised rather than being formally analysed using a structured approach (e.g. thematic analysis), will be excluded.
- Evaluation: Inductive themes representative of MCPs views and experiences of EWS use in clinical practice
- Research type: Published and unpublished studies reporting qualitative data on MCPs views or experiences of EWS application or use.

Search & selection methods

We will search the electronic databases systematically to identify and retrieve primary research studies. The databases that will be searched are MEDLINE, CINHAL, Web of Science Core Collection (incorporating Social Science Citation Index) and Maternity and Infant Care (MIDIRS). Databases will be searched from the date of inception to the date the search is implemented. Search terms were developed using elements of the SPIDER acronym (Table 1) and adapted as appropriate for the different databases. Language filters will not be applied to the search, although we will only include studies published in English. The rationale for searching all languages unfiltered is to identify potentially eligible non-English publications, and, depending on how many might be retrieved, whether possible language bias could be introduced by their exclusion. Records retrieved from the databases will be downloaded to Endnote (version EN20). Duplicate records will be removed and the remaining records uploaded to Covidence software for eligibility screening. Screening records against the review’s eligibility criteria will be undertaken by two members of the review team (VS & KC). All records will be initially screened based on their title and abstract. Those that appear eligible at this level will be forwarded for full text review. Following a review of full-text papers, studies meeting our inclusion criteria will be included in the review. Any uncertainty or disagreements regarding the inclusion or exclusion
of records will be resolved through discussion, or, if needed, the record will be assessed by a third review author (DOM) until a consensus decision is achieved.

We will expand our search by additionally searching the reference lists of studies identified for inclusion in the review and by searching the proceedings of international maternity care conferences (e.g. International Confederation of Midwives Triennial Conference 2017; Normal Labour and Birth Research Conference 2020). We will perform searches of grey literature databases (e.g. https://www.greylit.org/) and will search the reference lists of any identified national clinical guidelines on maternity EWS for potentially relevant studies that might not have been captured in our electronic database search.

Quality appraisal of included studies
Assessing the methodological quality of included studies formally is a key component of systematic reviews\(^27\), and perhaps even more-so in qualitative evidence synthesis as the process of assessment itself can facilitate a deeper understanding of the included studies\(^30\). A variety of appraisal tools are available for assessing the methodological quality of qualitative studies although these can range from having very broad criteria to consisting of explicit checklists\(^29\). For purposes of quality appraisal in this review we have chosen the appraisal tool developed and previously used\(^30\) by the Evidence for Policy and Practice Information and Co-ordinating (EPPI) Centre. The tool consists of 12 quality assessment criteria (A-L) across three core domains that are centred on study reporting, data collection and analysis, and study methods (Table 2). Two pairs of reviewers (VS & DOM; VS & KC) will independently assess the extent to which each quality criterion is met in each study. Irrespective of how many or how few quality criteria are met, we will include all studies for data extraction and synthesis purposes as qualitative studies of poor methodological may still provide important views data that could have considerable relevance to

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### Table 1. Search terms.

<table>
<thead>
<tr>
<th>S</th>
<th>Nurse* OR midwi* OR (public health nurse*) OR (practice nurse*) OR doctor* OR (general practitioner*) OR obstetrician* OR clinician OR physician OR (healthcare provider) OR (healthcare professional) AND</th>
</tr>
</thead>
<tbody>
<tr>
<td>PI</td>
<td>(Physiological track and trigger*) OR (early warning*) OR (early warning NEXT-1 (score OR tool, OR signs OR triggers OR system)) OR mews OR meows OR moews OR (escalation adj protocol*) (escalation adj policy) AND Maternity (MeSH) OR Obstetrics (MeSH) OR Pregnant* OR Pregnancy (MeSH) OR (antenatal OR prenatal OR perinatal OR puerperal OR puerperium OR postnatal OR postpartum OR peripartum OR post-natal OR post-partum OR ante-natal OR ante-partum OR obstetric*).tw OR Postpartum period (MeSH) AND</td>
</tr>
<tr>
<td>E or R</td>
<td>experiences OR experience OR view* OR perceptions OR perception OR voices OR narratives OR qualitative OR (mixed AD) method OR (grounded theory) OR phenomenology OR (action research)</td>
</tr>
</tbody>
</table>

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### Table 2. Quality appraisal criteria\(^30\).

<table>
<thead>
<tr>
<th>Quality of the study reporting</th>
<th>A= Aims and objectives clearly reported</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>B= Adequately described the context of the research</td>
</tr>
<tr>
<td></td>
<td>C= Adequately described the sample and sampling methods</td>
</tr>
<tr>
<td></td>
<td>D= Adequately described the data collection methods</td>
</tr>
<tr>
<td></td>
<td>E= Adequately described the data analysis methods</td>
</tr>
</tbody>
</table>

| There was good or some attempt to establish the | F= Reliability of the data collection tools |
|                                               | G= Validity of the data collection tools |
|                                               | H= Reliability of the data analysis |
|                                               | I= Validity of the data analysis |

| Quality of the methods | J= Used the appropriate data collection methods to allow for expression of views |
|                       | K= Used the appropriate methods for ensuring the analysis was grounded in the views |
|                       | L= Actively involved the participants in the design and conduct of the study |

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our synthesis. As Thorne advises, eliminating a study based on its fit with particular quality appraisal guidelines “may obscure a germ of possibility that, if used to interrogate the reports of other studies, could have led to important new angles of consideration”33, p7.

Data extraction and synthesis
Data extraction will be based on the aim of the review and will involve extracting the following information:

- Year the study was published
- Aim of the study
- Funding details (if provided)
- Description of the participants and the study setting
- Study duration/timeframes
- Description of the maternity EWS (where provided)
- Method(s) of data collection and analysis
- Findings related to providers’ views and experiences of EWS application and use in maternity care

Using a standardised data extraction form35, the data will be extracted independently by two reviewers followed by accuracy cross-checks. Thematic synthesis, as described by Thomas and Harden for synthesising data from qualitative studies34 will be used to synthesise the studies’ data. In using this method, the following will be undertaken:

i) Manual line by line coding of extracted data; the extracted text including relevant participant quotes will be reviewed and coded by one member of the review team.

ii) Descriptive themes will be identified by assessing similarities and differences between codes which will be clustered into descriptive themes. Although one member of the review team will identify the descriptive themes, the review team will meet to discuss these themes to ensure they are reflective of the codes before progressing the synthesis to developing the analytical themes.

iii) Generate the analytical (or dominant) themes and sub-themes from the descriptive themes. One member of the review team will generate the analytical themes, however, all members of the review team will be involved in a process of reflection, iteration, and discussion in determining the final themes to ensure they are collectively representative of the studies’ data. This process will enhance synthesis rigour and transparency.

Assessment of confidence in the review findings; GRADE-CERQual
Grading of Recommendations Assessment, Development and Evaluation-Confidence in the Evidence from Reviews of Qualitative research (GRADE-CERQual)35–40 will be used to assess levels of confidence in the review findings. Using GRADE-CERQual, distinct review findings are assessed on the components of methodological limitations, coherence, extant or adequacy of contributing data and relevancy to the review question. An overall assessment of High, Moderate, Low or Very Low confidence in each finding, based on the component ratings, is then made36. For purposes of the assessments, we have established a priori downgrading criteria as illustrated in Table 3. Judgements are based on an initial assumption of ‘High confidence’ in all findings, and then downgraded accordingly. GRADE-CERQual assessments will be performed by two reviewers independently, as recommended35, with overall confidence levels based on discussions and consensus.

Table 3. GRADE-CERQual downgrading criteria.

<table>
<thead>
<tr>
<th>GRADE</th>
<th>Downgrading criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>- No or minor concerns in all four components</td>
</tr>
<tr>
<td></td>
<td>- Moderate concerns in one component and no concerns in remaining three, or minor concerns in one other component</td>
</tr>
<tr>
<td>Moderate</td>
<td>- Moderate concerns in one component, minor concerns in two components, and no concerns in remaining component</td>
</tr>
<tr>
<td></td>
<td>- Moderate concerns in two components and no concerns in remaining two components</td>
</tr>
<tr>
<td>Low</td>
<td>- Moderate concerns in two components and minor concerns in at least one other component</td>
</tr>
<tr>
<td></td>
<td>- Moderate concerns in three components and no concerns in remaining component</td>
</tr>
<tr>
<td></td>
<td>- Severe concerns in one component and no concerns in remaining three components, or minor concerns in one other component</td>
</tr>
<tr>
<td>Very Low</td>
<td>- Moderate concerns in three components and minor concerns in remaining component</td>
</tr>
<tr>
<td></td>
<td>- Moderate concerns in all four components</td>
</tr>
<tr>
<td></td>
<td>- Severe concerns in one component and moderate concerns in at least one other component</td>
</tr>
</tbody>
</table>
facilitators for EWS in practice based on the experiences of those directly involved in using EWS as part of maternity care provision.

In disseminating the findings of the QES, the target audience will be primarily maternity care providers and local or national guideline or policy developers. For this reason, we aim to publish the findings in a healthcare journal that predominantly publishes maternity care research and is known to have a wide maternity care provider readership. Summary findings in the form of short reports will be prepared and shared with national guideline groups (e.g. National Clinical Effectiveness Committee, Ireland), and will be made available to practice development departments in maternity settings. Links to the published report will be shared on social media (Twitter and Facebook), and summary findings distributed via online maternity/midwifery email lists, as appropriate.

Review status
The search strategy has been implemented. Screening (level I, title and abstract) the retrieved records for eligibility is currently in progress.

Data availability
Underlying data
No data are associated with this article.

Extended data

This project contains the following extended data:
- Template Data Extraction Form.docx

Reporting Guidelines

Data are available under the terms of the Creative Commons Attribution 4.0 International license (CC-BY 4.0).

References


Open Peer Review

Current Peer Review Status: ✔ ✔

Version 1

Reviewer Report 05 January 2022

https://doi.org/10.21956/hrbopenres.14443.r31044

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Mirjam Lukasse

1 Department of Nursing and Health Promotion, Oslo Metropolitan University, Oslo, Norway
2 Department of Health and Social Sciences, University of South-Eastern Norway, Vestfold, Norway

This is a study protocol for a qualitative evidence synthesis of maternity care providers' views on and experiences with a type of screening tool to identify a deterioration in a patient's physical condition as early as possible, called an Early Warning System (EWS).

To date, a synthesis of the qualitative evidence has not been published while there are a number of original research papers in this field. So I am pleased these researchers are willing to take on the work to perform this synthesis. I find the proposal well written. The authors show they know the field well. The proposed methodology is solid. I look forward to reading the synthesis.

I have just a couple of minor questions. In the methods part of the abstract it is written that studies will be included from the date of inception. Quite a few readers will not know when this was, and this date may also vary from country to country. Could this phrase be replaced by a year that ensures all studies are included?

The authors will only include studies published in English, but report on how many they find in other languages. There may of course not be any papers in other languages. However, I would like the authors to consider having the papers translated if they find one of two in another language. Good information may be lost by not doing this. I realize that English is the dominant language for publication of research. However, midwives who may investigate the introduction of an early warning system in their practice environment may not be such advanced researchers, but can still produce good research under supervision while doing, for example, a Master in Midwifery. If such a study is published in French or German it would be very sad not to have it included. However, this is just a suggestion.

Is the rationale for, and objectives of, the study clearly described?
Yes

Is the study design appropriate for the research question?
Yes

Are sufficient details of the methods provided to allow replication by others?
Yes

Are the datasets clearly presented in a useable and accessible format?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Midwifery and midwifery education

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 19 October 2021

https://doi.org/10.21956/hrbopenres.14443.r30302

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Ruth Endacott
MEDICINE, NURSING AND HEALTH SCIENCES, Monash University, Melbourne, VIC, Australia

The proposed qualitative evidence addresses an important, timely, and under-reviewed area of clinical practice - experience of maternity care providers in the use of Early Warning Systems (EWS). Existing evidence in non-maternity health systems emphasises the impact of human factors on decisions about use of EWS and implementation of escalation. This is a comprehensive, appropriately referenced, protocol for a qualitative evidence synthesis. The rationale is clear and review processes are mostly transparent. A few minor changes would make the methods more transparent and replicable for future researchers:

1. Some of the evidence might lie in medical education literature, for example use of EWS in simulated situations. EWS is a key part of education for obstetric emergencies and some studies include qualitative interviews or photo elicitation interviews so might be a valuable source. It is not clear whether these studies are excluded.

2. Under Methods, it would be helpful to use 'eligibility criteria' as the sub-heading, and then address inclusion criteria and exclusion criteria (i.e. when you have applied your inclusion criteria, what studies would you then exclude?).

3. A word is missing in the following sentence: we will include all studies for data extraction and synthesis purposes as qualitative studies of poor methodological may still provide...

4. It would be helpful to add a sentence or two to outline how you will link the thematic synthesis
and the output from the GRADECERQual process. This is important for replication of your method but also to ensure that clinicians can easily apply (or not) the QES findings to clinical practice, education of maternity care providers and future policy guidance for EWS in maternity services.

**Is the rationale for, and objectives of, the study clearly described?**
Yes

**Is the study design appropriate for the research question?**
Yes

**Are sufficient details of the methods provided to allow replication by others?**
Partly

**Are the datasets clearly presented in a useable and accessible format?**
Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** managing deterioration

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.