ABSTRACT

Introduction: Studies suggest that continuous glucose monitors (CGMs) play an important role in the management of diabetes. Although general acceptance has been reported by patients with type 2 diabetes towards the use of CGMs, potential barriers exist like pain due to sensor insertion, accidental removal of the device or adhesive strip, impacts of daily activities, skin reactions to sensor adhesive, etc. This systematic review of qualitative studies aims to explore the perspectives, experiences and narratives of patients and caregivers about CGM use, and its barriers and facilitators.

Methods and analysis: This review will include qualitative studies and cross-sectional and longitudinal cohort studies using open-ended questions, published in English by 30 October 2021. The following electronic databases will be searched: Cochrane Library, PubMed, EMBASE, CINAHL, PsycINFO and Scopus. A search of grey literature will be conducted via an online search of Google Scholar, WorldCat, ClinicalTrials.gov and OpenGrey. A combined search strategy using medical subject headings (MeSH); controlled vocabulary and ‘free-text’ terms will be appropriately revised to suit each database. Primary outcomes will include patient and caregiver perspectives on diabetes management regarding glucose control; living with CGM (quality of life, experience of wearing a CGM); psychological aspects (anxiety, depression, emotional burden); barriers (technical issues, financial issues) to use of CGM and thoughts (interpretation, understanding) on the CGM report. A qualitative meta-synthesis will be conducted employing a systematic literature search of existing literature, quality assessment using study-specific tools and an aggregative thematic synthesis by a multidisciplinary team.

Ethics and dissemination: Ethical approval is not required since this is a systematic review. The results will help improve clinical implementation of CGMs on part of both patients and caregivers.

PROSPERO registration number: CRD42020152211.

INTRODUCTION

The prevalence of diabetes mellitus is predicted to increase to 10.9% (700 million) by 2045. The costs due to diabetes mellitus and its complications are also predicted to increase up to $2.1 trillion by 2030. Globally, around 1 in 11 adults have diabetes mellitus out of which 90% have type 2 diabetes mellitus (T2D). Therefore, it is essential to explore effective methods of diabetes management to alleviate the global health burden caused by T2D and its complications.

Continuous glucose monitoring (CGM) is a device that can evaluate the blood glucose fluctuations in real time and can help develop personalised treatment plans to fully control short-term fluctuations in blood glucose levels. Many studies have shown its potential in diabetes management through...
identification of hyperglycaemia and hypoglycaemia, measuring glycaemic control and providing actionable information in the form of the CGM report to health carers and patients. Studies have also reported general acceptability by patients with T2D. It is estimated that real-time CGM can lead to higher quality-adjusted life years and reduced healthcare costs, with an expected incremental cost-effectiveness ratio of €180,553 per quality-adjusted life year for patients with T2D. Therefore, CGM might be an effective method for T2D management. Even though the use of CGMs in adults with T2D is recommended by the endocrine society clinical practice guidelines, its uptake is less among people with T2D compared with people with type 1 diabetes (T1D).

Research amongst using CGMs has identified several barriers to its use. A qualitative review exploring the impact and experience of using CGMs among people with T1D reported that CGM affects physical, emotional and relational aspects of life. It also found that clinicians can provide education and management to help reduce the barriers of CGM use. In 2018, a systematic appraisal of personal blogs analysing 39 blogs found that real-time CGM data sharing enhanced the feelings of safety among patients with T1D and their health care providers. A recent review reported pain associated with insertion and wear of CGMs, body image issues, alarm fatigue, information overload, accuracy concerns, and clinical inertia to be barriers against CGM use among patients with T1D. CGMs have been widely used in the management of T1D, but their effectiveness has led to their use for management of T2D. Therefore, research needs to extend to exploration of experiences, barriers and facilitators to use of CGM among patients with T2D.

Effectiveness of CGM in T2D management has been reported through quantitative studies. However, only a few qualitative studies have explored its feasibility. In order to understand its wider acceptability, it is important to explore the narratives of people using CGM for management of T2D and their caregivers; and gain understanding of facilitators and barriers from the perspectives of both patients and caregivers. To improve its usage for clinical or family settings, CGM device functions are improved regarding user satisfaction and usability over time. It is suggested that use of CGM should consider the users’ preferences and their assessment of acceptance. Therefore, both the experiences and thoughts of caregivers and users should be explored and generated among population with T2D.

Although general acceptance has been reported by patients with T2D towards the use of CGMs, potential barriers like pain due to sensor insertion, accidental removal of the device or the adhesive strip, impacts of sports and daily activities, skin reactions to sensor adhesive, lack of insurance coverage, etc exist. These barriers have been identified by qualitative studies, that also provide valuable perspectives for promoting the use of CGMs for clinical and self-management of diabetes. The studies also identified a gap in literature and a need for a meta-synthesis of qualitative studies on this topic. Therefore, this systematic review aims to explore the perspectives, lived experiences and narratives of patients and caregivers about their CGM use, and its barriers and facilitators.

METHODS AND ANALYSIS

Research questions

This qualitative meta-synthesis aims to answer the following research questions:

1. How do people with T2D describe their experience (ie, glucose control, living with CGM, psychological aspects, understanding of CGM reports) of using CGM for managing their diabetes?
2. What are the views of people with T2D who use CGM for managing their diabetes about its effectiveness in maintaining glucose control?
3. What are the barriers and facilitators to CGM use reported by people with T2D and their caregivers; and what can the healthcare providers (ie, doctors, nurses, dietitians) do to address these barriers?

Design and eligibility

A qualitative meta-synthesis will be conducted for existing published and unpublished literature following a systematic literature search, quality assessment using study-specific tools, aggregative thematic synthesis and the reporting of analytical themes to highlight the research questions in accordance with the guidelines of the Enhancing transparency in reporting the synthesis of qualitative research (ENTREQ). Preferred Reporting Items for Systematic Review and Meta-Analysis Protocols (PRISMA-P) (online supplemental file 1) was used to prepare this protocol.

This appropriate structure of review is designed based on the Joanna Briggs Institute (JBI) template. It will include published and unpublished qualitative studies irrespective of the study design and cross-sectional and longitudinal cohort studies using open-ended questions that report perspectives and experiences of using CGM of patients with T2D and their caregivers. We will exclude studies of secondary designs (after checking the list of included studies for potential relevant primary studies), and studies only using quantitative methods. The inclusion of studies is primarily qualitative (ie, interviews, semi-structured interviews, focus group discussions) studies and mixed-methods research (ie, questionnaires that are qualitative in nature with open-ended questions). The full list of inclusion and exclusion criteria has been presented in table 1.

Information sources and search strategy

A comprehensive PubMed search strategy was formulated after consultation with an experienced medical librarian (online supplemental file 2). Comprehensive search strategies for other databases will be developed using medical subject headings (MeSH), controlled vocabulary
The study selection will be done with Covidence, and study selection and management disagreements will be resolved by a third reviewer (AK) through discussion. A PRISMA flow diagram will be formed to record the details of the study selection process (online supplemental file 3).

Data on study characteristics will be extracted into an adopted JBI data extraction form by two reviewers (MZ and AP) and will include first author, year of publication, country, research questions, study population (age, ethnicity, gender, socioeconomic status), number of participants, study setting, data collection method, data analysis method, themes relevant to the objectives of this review and study limitations. Authors will be contacted via email to obtain relevant additional or missing information if required.

The analysis of included studies will follow the Standards for Reporting Qualitative Research: A Synthesis of Recommendations with 21 items. The thematic framework analysis approach will be adopted for data synthesis and analysis. The research team will follow the six stages of framework synthesis using NVivo (V.12.0.0 Plus): (a) procedure for analysis; (b) familiarisation with the interview; (c) coding; (d) developing a working analytical framework; (e) applying the analytical framework and (f) charting data into the framework matrix. Initial coding of a sample set (20%) of papers will be conducted by two reviewers (MZ and AP) independently and in consultation with a third reviewer (AK) to agree on the outline of the coding framework. Following this, coding of the remaining studies will be undertaken in duplicate with regular discussion to review and agree on additions and amendments to the framework.

The extracted data will be analysed line by line, and the identity of each first-order or second-order structure will be coded accordingly. The code will be inductively created by two independent reviewers for the response to uncovered findings, with a third reviewer’s consultation. The two reviewers (MZ and AP) will then organise an expert meeting with team members to finalise the theme names and the structure/hierarchy of codes/themes. The thematic analysis will be included but not limited to the following categories (online supplemental file 4), the final themes will be decided after the expert meeting based on the evidence available as it will follow

Supplemental File 3).


## Table 1 Inclusion and exclusion criteria

<table>
<thead>
<tr>
<th>Concepts</th>
<th>Inclusion</th>
<th>Exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Type of study*</td>
<td>Primary qualitative (ie, interviews, semistructured interviews, focus group discussion) studies</td>
<td>Quantitative studies without qualitative component, conference reports, commentaries, opinion pieces, business reports, case reports, abstracts and systematic reviews. Studies adopting mix-methods but only reporting quantitative data</td>
</tr>
<tr>
<td>Study population</td>
<td>Age ≥18 years; a confirmed diagnosis of T2D; wearing any types of CGMs for at least 3 days</td>
<td>Adolescents (under 18 years of age) and children; other types of diabetes (ie, T1D, gestational diabetes mellitus)</td>
</tr>
<tr>
<td>Language</td>
<td>English</td>
<td>Non-English</td>
</tr>
<tr>
<td>CGM wearing</td>
<td>Patients wearing any type of CGMs (ie, real-time CGM, retrospective CGM, flash CGM) for at least 3 days</td>
<td>Patients wearing CGMs for less than 3 days</td>
</tr>
</tbody>
</table>

*Inclusion will not be limited to designs such as phenomenological, grounded theory, ethnography, action and feminist research.

CGM, continuous glucose monitor; T1D, type 1 diabetes; T2D, type 2 diabetes.

and ‘free-text’ words relevant to this review, and the logic grid will be developed with the research team (online supplemental file 2). Reference lists of relevant systematic reviews will be hand-searched to identify additional relevant studies or reviews. Topic experts in the field will also be contacted to identify any unpublished or ongoing work. Additional literature will be sought by checking the reference lists of journal and conference articles, and checking cited articles. The research team will re-run the searches before the final systematic review will be published to ensure that they have not missed the latest studies.

The search strategy will aim to find both published and unpublished studies in English language published up to 30 October 2021 from the following databases: Cochrane Library, PubMed, EMBASE, CINAHL, PsycINFO and Scopus. To find unpublished grey literature (ie, theses and dissertations), Google Scholar, WorldCat, Clinical-Trials.gov and OpenGrey will be searched.

### Data management

The reference management system EndNote V.X9 will be used to manage studies exported from all the sources. Two reviewers (MZ and AP) will independently and in duplicate screen the records for inclusion at the title/abstract and full-text level screening via Covidence online system (https://www.covidence.org). Any disagreements at any screening level will be resolved through discussion. For any unresolved decisions, a third reviewer (AK) shall be consulted.

### Study selection and management

The study selection will be done with Covidence, and includes title and abstract screening and full-text screening and extraction. Disagreements will be resolved by third and fourth reviewers (AK and YL) through discussion. A PRISMA flow diagram will be formed to record the details of the study selection process (online supplemental file 3).

Data on study characteristics will be extracted into an adopted JBI data extraction form by two reviewers (MZ and AP) and will include first author, year of publication, country, research questions, study population (age, ethnicity, gender, socioeconomic status), number of participants, study setting, data collection method, data analysis method, themes relevant to the objectives of this review and study limitations. Authors will be contacted via email to obtain relevant additional or missing information if required.

The analysis of included studies will follow the Standards for Reporting Qualitative Research: A Synthesis of Recommendations with 21 items. The thematic framework analysis approach will be adopted for data synthesis and analysis. The research team will follow the six stages of framework synthesis using NVivo (V.12.0.0 Plus): (a) procedure for analysis; (b) familiarisation with the interview; (c) coding; (d) developing a working analytical framework; (e) applying the analytical framework and (f) charting data into the framework matrix. Initial coding of a sample set (20%) of papers will be conducted by two reviewers (MZ and AP) independently and in consultation with a third reviewer (AK) to agree on the outline of the coding framework. Following this, coding of the remaining studies will be undertaken in duplicate with regular discussion to review and agree on additions and amendments to the framework.

The extracted data will be analysed line by line, and the identity of each first-order or second-order structure will be coded accordingly. The code will be inductively created by two independent reviewers for the response to uncovered findings, with a third reviewer’s consultation. The two reviewers (MZ and AP) will then organise an expert meeting with team members to finalise the theme names and the structure/hierarchy of codes/themes. The thematic analysis will be included but not limited to the following categories (online supplemental file 4), the final themes will be decided after the expert meeting based on the evidence available as it will follow
an inductive analysis approach. The goal of this phase is to go beyond the initial findings to generate more understanding and information for future studies.

Quality assessment
Critical Appraisal Skills Programme (CASP) tool for mixed-methods studies (https://casp-uk.net/casp-tools-checklists/) will be used to critically appraise the quality of included studies using mixed-methods. This 10-item CASP tool is considered to be the most suitable tool to consider the quality parameters of qualitative work, and is a well-accepted and validated tool.26 JBI Critical Appraisal Checklist for Qualitative Research will be used to critically appraise the quality of included qualitative studies.27 Two independent reviewers (MZ and AP) will conduct the critical appraisal independently and in duplicate of all studies included for full text. If required, a third reviewer (AK) will identify the differences during risk of quality assessment. Studies will not be excluded based on their quality rating unless there are serious concerns (ie, very poor quality due to lack of transparency in reporting of methods or lack of reporting of participant quotations).

Data synthesis and thematic analysis
The strategy for the data synthesis of this review is based on the System for the Unified Management, Assessment and Review of Information (SUMARI) with the meta-aggregation approach.28 A preliminary and comprehensive synthesis will be formed for the following aspects:29 thematic and content analysis. The narrative synthesis of the review, tables and narrative summaries of each included study will be illustrated by ATLAS.ti, data analysis software, following the Guidance of Narrative Synthesis in Systematic Reviews.25 This will involve a summary or synthesis of discoveries to concatenate the discoveries by combining the discoveries and categorising them based on similarities to generate a set of statements representing the set. These categories will then be subjected to a synthesis in order to produce a single comprehensive set of synthesised findings that can be used as a basis for evidence-based practice. Where textual pooling is not possible, the findings will be presented in narrative form.

Primary outcomes will include patient and caregiver perspectives on diabetes management regarding glucose control; living with CGM (quality of life, experience of wearing CGM devices); psychological aspect (anxiety, depression, emotional burden); barriers (technical issues, financial issues) of CGM use and thoughts (interpretation, understanding) on the CGM report. The outcomes will be modified depending on the evidence available. Secondary outcomes will include weight management, change in blood pressure and lipid profile.

To consider the impact on different systems, we will categorise different types of CGMs for additional analysis: (1) real-time CGM, (2) professional CGM and (3) flash/intermittently scanned CGM. In our report, we will tabulate summaries of the characteristics of the included studies. Important limitations and gaps within the evidence base will be presented and discussed.

Patient and public involvement
Patients and the general public will not be involved directly in the design and conduct of this review. However, the development of the review questions was informed by patient safety concerns and the experience of health professionals using eHealth applications in clinical practice. All data analysed during this study are included in the manuscript and additional files.

Ethics and dissemination
Ethical approval is not required as this systematic review will use secondary data. This systematic review protocol is registered in the International Prospective Register of Systematic Reviews (http://www.crd.york.ac.uk/PROSPERO). The results will help improve clinical implementation of CGMs for both patients and caregivers. This systematic review will be published in a peer-reviewed journal and presented in scientific conferences.

DISCUSSION
To the best of our knowledge, this is the first systematic review synthesising qualitative evidence exploring barriers and facilitators to CGM use among patients with T2D and their caregivers through exploration of their narratives of their perspectives and experiences. To ensure reliability, robustness and transparency of the predefined methods to answer the research questions of this review, we have referred to the SUMARI, JBI and ENTREQ guidelines.21 27 28 Thematic analysis can identify prominent topics and process the literature on these topics in an organised and structured manner. It is flexible, allows reviewers considerable freedom, and is a method of integrating qualitative and quantitative evidence.30 Moreover, thematic analysis can be either theory driven (oriented to the appraisal of specific themes by interrogation of the study) or data driven (driven by the themes identified in the study itself).30 This review will adopt the data-driven thematic analysis to explore and analyse25 patients’ and caregivers’ experiences.

The results will be made available through publication in a peer-reviewed journal and through local and international presentations. This study will help identify barriers and facilitators to CGM use from the existing literature and provide scientific evidence for future studies regarding clinical use of CGMs for patients, caregivers and healthcare providers. This review will also provide future directions for observational and experimental studies regarding T2D management by avoiding issues resulting from the use of different types of CGMs. The findings of this qualitative evidence synthesis may help to explain why and how some interventions to improve CGM uptake.
are more effective than others. Knowledge of perceived facilitators and barriers to CGM use may generate insights about potential causal factors that affect impact. By identifying influential factors, it may also contribute to the development of more relevant, acceptable and effective interventions in the future. The findings can also be used to inform the design of future effectiveness reviews suggesting outcomes that are important to patients with T2D and their caregivers, as well as generating hypotheses that can be tested out, for example, in future subgroup analyses. In addition, the results from this synthesis may help improve our understanding of the reasons for acceptance of CGMs from the perspective of patients with T2D and their caregivers, contributing to the future development of more relevant, acceptable, and in turn, effective interventions to promote patient acceptance and uptake of CGMs.

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Contributors
MZ and PH conceived and designed the study. MZ drafted the manuscript and is the guarantor of the systematic review. AP developed the research questions and study design. MZ and YL designed the tables for included studies and will evaluate the quality of included studies in the systematic review. AP, MZ, YL, AK and XZ revised the manuscript. All authors reviewed and approved the final manuscript as submitted and agreed to be responsible for all aspects of the work.

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Competing interests
None declared.

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Provenance and peer review
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Supplemental material
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