

# Healthcare Providers' Implementation of Patient-Centred Outcome Measures (PROMs) and Patient-Reported Experience Measures (PREMs) in Practice A Protocol for a Mixed Method Systematic Review

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## Protocol

**Keywords:** Patient reported outcome measures (PROMs), Patient reported outcomes (PROs), Routine outcome monitoring (ROM), Patient reported experience measures (PREMs), Knowledge translation (KT), Patient-centred care (PCC), Clinical decision-making, Clinicians, Healthcare providers, Implementation

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## Abstract

**Background:** Substantial literature has highlighted the importance of patient-reported outcome and experience measures (PROMs and PREMs, respectively) to collect clinically relevant information to better understand and address what matters to patients. The purpose of this systematic review is to synthesize the evidence about strategies for implementing individual-level PROMs and PREMs data by healthcare providers in their everyday practice.

**Methods:** This mixed method systematic review protocol describes the design of our synthesis of the peer-reviewed research evidence (i.e., qualitative, quantitative, and mixed methods), systematic reviews, organizational implementation projects, expert opinion, and grey literature. Keyword synonyms for “PROMs,” “PREMs,” and “implementation” will be used to search eight databases with limiters of English from 2009 to 2019. Study selection criteria includes implementation at the point-of-care by healthcare providers in any practice setting. Eligible studies will be critically appraised using validated tools (e.g., Joanna Briggs Institute). Guided by the review questions, data extraction and synthesis will occur simultaneously to identify biographical information and methodological characteristics as well as classify study findings related to implementation processes and strategies. As part of the narrative synthesis approach, two frameworks will be utilized: (a) Consolidated Framework for Implementation Research (CFIR) to identify influential factors of PROMs and PREMs implementation and (b) Expert Recommendations for Implementing Change (ERIC) to illicit strategies. Data management will be undertaken using NVivo 12™.

**Discussion:** Data from PROMs and PREMs are critical to adopt a person-centred approach to healthcare. Findings from this review will guide subsequent phases of a larger project that includes interviews and a consensus-building forum with end users to create guidelines for implementing PROMs and PREMs at the point of care.

**Systematic Review Registration:** PROSPERO CRD42020182904

## Background

Shifting from traditional, biomedical, disease-focused, and scientific models of healthcare toward patients being drivers of their care is essential to improve health and clinical outcomes (1). Person-centred care that focuses on what matters to patients occurs when healthcare providers (HCPs) include and listen to the voice of patients. To do this, data collected from patient-reported outcome and experience measure (PROMs and PREMs, respectively) can be used by HCPs to incorporate patients’ voice and inform their care (2). Our aim is to conduct a mixed methods systematic review to synthesize existing knowledge on effective methods for implementing individual-level PROMs and PREMs data by HCP in their everyday practice. The protocol for this systematic review is part of a larger study to create user guidelines that support clinical decision-making and care planning, wherein HCP interviews and a deliberative dialogue, consensus-building forum with key stakeholders will also contribute to guideline development. The patient-centred measurement (PCM) implementation guidelines for HCPs will support routine collection, interpretation, and integration of these data in everyday practice, ultimately contributing to effective, quality health care.

PCM is an umbrella term used in this systematic review to include tools for measuring patients’ experiences or outcomes (3). Substantial literature has highlighted the importance of PCM measures to collect clinically relevant information from patients to better understand and address what matters to them. PROMs are questionnaires (also referred to as assessment tools or instruments) to measure patients’ health status in a standardized and quantitative manner at a point in time (3). These inquiries may capture how patients function or feel with respect to their health; disease condition and its treatment; or functional status, quality of life, or mental wellbeing (4, 5). In contrast, PREMs focus on how patients feel with respect to their health care or illness experience. PREMs are commonly used to solicit information about patient satisfaction with service delivery in a clinical setting, or describe patient experience of a therapy or plan of care (3). To inform and plan patients’ care, PROMs and PREMs can be collected by HCPs for various purposes: screening, assessment, monitoring, planning, and evaluating care (e.g., interventions, treatment, referrals, and tests) as well as creating decision aids (6, 7). Both PROMs and PREMs are intended to provide assessment data about patients’ health thereby complimenting HCPs’ use of clinician-based outcomes (CBOs), biological measures, and physical examination. When developed with input from patients and regularly integrated into clinical practice, PROMs and PREMs data can encourage conversations between patients and HCPs. As a result, such conversations lead to shared decision-making, improved patient-clinician communication, detection of overlooked problems, and tailored process monitoring thereby ensuring quality individualized care (4, 8-10).

The use of PROM and PREM data in healthcare has been a robust area of research at individual, organizational, and system levels around the world. Considerable emphasis has been placed on the use of aggregated PCM data to inform program evaluation, quality improvement, benchmarking, value-based healthcare, and to some degree managerial decision making (11, 12). Over several decades, developers and users have examined PROM and PREM applications to clinical practice (e.g., (6, 10, 13-16)). Although structures and processes exist to support the use of aggregated PCM data, the routine integration of individual-level PCM data by HCPs in clinical practice is challenging worldwide for various reasons, with calls for additional research to understand best practices for effective implementation strategies with a focus on end users (3, 17-20). Eight systematic reviews (5, 10, 11, 21-25) and one scoping review (26) have explored PCM implementation by HCPs (see Table 1). While this number of systematic reviews would typically offer a robust body of evidence, a closer inspection reveals a paucity of literature on effective knowledge translation strategies for PCM implementation by HCPs at the point of care. Five reviews identified barriers and facilitators of PCM use as one of several aims (5, 10, 23, 25, 26), with only three stating this as the primary topic often in a specific area of practice (e.g., oncology) (11, 22, 24). Notably, Bantug and colleagues (11) differed from other reviewers in their examination of effective methods for the interpretation of patient-reported data using various graphic displays. Our systematic review will synthesize existing evidence about the everyday use of individual-level PCM data by HCPs in all practice settings or healthcare context.

Table 1. Summary of Published Systematic and Scoping Reviews about Using PCM

Author Name in Chronological Order	Objectives	Phenomenon of Interest Pertinent to this Review And Clinical Area	Types of Evidence Included and Synthesis Method	Databases Searched and Search Dates	Search Terms for Patient-Reported Measures	Included # Articles (Search Results)	Inclusion Criteria Pertaining to HCPs	Quality or Bias Assessment Conducted
Duncan 2012 <sup>2</sup> (32)	What are the barriers and facilitators to routine outcome measurement by allied health professionals in practice? (p. 2)	Barriers and facilitators to routine outcome measurement  Area: Any setting	No restriction on study design  Narrative synthesis	MEDLINE, CINAHL, PsycINFO  ≤ 2010	13 synonyms	15 (960)	Concerned with identifying or researching factors which acted as facilitators and/or barriers in the routine use of outcome measures by allied health professionals in practice.	Yes
Antunes, 2014 <sup>1</sup> (5)	1. Identify barriers and facilitators to the systematic implementation of PROMs in palliative care clinical practice. 2. Identify needs and other comments of clinical teams regarding the routine use of PROMs. 3. Identify lessons learned on the process of implementation of PROMs in clinical practice (p. 160).	Barriers and facilitators to implementation of PROMs  Lessons learning on the PROMs implementation process  Area: Palliative	All study designs  Narrative synthesis	MEDLINE, CINAHL, PsycINFO, Embase, British Nursing Index  1985-Mar 2011	Example for British Nursing Index  Outcome measure* PROs PROMs Patient reported outcome? Patient reported outcome measure? Self-reported measure? (Appendix 1)	31 (3863)	Not stated	Yes
Boyce 2014 <sup>1</sup> (11)	Investigated the experiences of healthcare professionals with the use of PROMs . . . and to synthesise findings about the barriers and facilitators to their use (p. 509).	Use of PROMs Barriers and facilitators of PROMs use  Area: Any setting	Qualitative  Thematic analysis	PubMed, CINAHL, PsycINFO  ≤ Aug 2013	12-15 synonyms depending on database	16 (8344)	Healthcare professionals. Professionals' views of PROMs after receiving PROMs feedback  about individual patients or groups of patients.	Yes

Author Name in Chronological Order	Objectives	Phenomenon of Interest Pertinent to this Review And Clinical Area	Types of Evidence Included and Synthesis Method	Databases Searched and Search Dates	Search Terms for Patient-Reported Measures	Included # Articles (Search Results)	Inclusion Criteria Pertaining to HCPs	Quality or Bias Assessment Conducted
Howell 2015 <sup>1</sup> (26)	<p>1. Which PROMs does the published, English literature show have been implemented for use in routine cancer clinical practice and in what phases of the trajectory?</p> <p>2. What are the barriers and enablers influencing clinical uptake of PROMs in routine care?</p> <p>3. What is impact of the routine use of PROMs on outcomes at the patient, provider, and system levels? (p. 1846-47)</p>	<p>Barriers and enablers influencing clinical update of PROMs</p> <p>Area: Cancer/oncology</p>	<p>Quantitative, qualitative, or systematic literature reviews</p> <p>Scoping review</p>	<p>MEDLINE, CINAHL, PsycINFO</p> <p>2003-2013</p>	<p>Examples given included</p> <p>self-report questionnaire</p> <p>self-assessment</p> <p>PRO</p> <p>PROMs</p> <p>PROMIS</p> <p>Outcomes</p>	30 (2447)	<p>Evaluated outcomes at the patient, clinical practice, or care process or system-level or barriers/</p> <p>enablers to the uptake or use of PROMs</p>	Not specified
Bantug 2016 <sup>1</sup> (21)	<p>Review existing literature examining the interpretation of PRO data displayed graphically (p. 484)</p>	<p>Interpretation of PRO data</p> <p>displayed graphically</p> <p>Area: Any setting</p>	<p>Quantitative or qualitative</p> <p>Integrative approach</p>	<p>MEDLINE</p> <p>1999-2014</p>	<p>Patient-reported outcome</p> <p>routine clinical care</p> <p>health-related quality of life</p> <p>health outcomes (p. 485)</p>	9 (242)	<p>Group- or individual-level PRO data for clinicians or patients</p>	Not specified
Foster 2018 <sup>1</sup> (24)	<p>To conduct a systematic review of reviews to identify the facilitators and barriers to implementing PROMs in organisations delivering health related services (p. 2-3)</p>	<p>Facilitators and barriers to impacting the implementation of PROMs</p> <p>Area: Any setting</p>	<p>Systematic reviews</p> <p>Framework synthesis</p>	<p>MEDLINE, CINAHL, PsycINFO, Embase, Cochrane Database of Systematic Reviews</p> <p>≤ 2017</p>	<p>PROM, PROMS, patient reported outcom*, PROs, PRO, PREM, PREMs, patient reported experience measure*, ePROs, ePRO, or outcome measure*</p>	6 (2047)	<p>Focus on health-related services irrespective of the type of provider.</p>	Yes

Author Name in Chronological Order	Objectives	Phenomenon of Interest Pertinent to this Review And Clinical Area	Types of Evidence Included and Synthesis Method	Databases Searched and Search Dates	Search Terms for Patient-Reported Measures	Included # Articles (Search Results)	Inclusion Criteria Pertaining to HCPs	Quality or Bias Assessment Conducted
Yang 2018 <sup>1</sup> (10)	<p>1. Who uses PROs as a tool to improve communication with patients and which topics are discussed?</p> <p>2. Through what mechanisms do PROs facilitate communication between clinicians and patients?</p> <p>3. What factors act as barriers or facilitators to PRO use and improved patient-clinician communication? (p. 42)</p>	<p>Barriers or facilitators to PRO use</p> <p>Area: Cancer/oncology</p>	<p>Not specified</p> <p>Realist synthesis</p>	<p>MEDLINE, CINAHL, PsycINFO, Embase, Cochrane Database of Systematic Reviews, Cab Direct</p> <p>≤ 2016</p>	<p>Example MEDLINE</p> <p>PROM</p> <p>PROMs</p> <p>PROMIS</p> <p>patient reported outcome*</p> <p>patient* outcome* measure*</p> <p>self report adj1 (questionnaire or measure)</p> <p>Exp *Health Status Indicators/ Questionnaires/ *Psychometrics/ Self Report/ Health Surveys/</p>	43 (610)	<p>PROs were used as an intervention, not as a study outcome</p> <p>measurement tool</p>	Yes
van Egdom 2019 <sup>1</sup> (23)	<p>1. How have PROMs been administered in breast cancer care?</p> <p>2. What is the impact of PROM administration on patients, care providers, and healthcare services or processes?</p> <p>3. What are the facilitators and barriers that influence the integration of PROM</p> <p>collection in routine breast cancer clinical practice? (p. 1198)</p>	<p>Facilitators and barriers that influence the integration of PROM</p> <p>Collection</p> <p>Area: Cancer/oncology</p>	<p>Excluded case report, editorial, review or study protocols</p> <p>Not specified</p>	<p>MEDLINE, CINAHL, PsycINFO, Embase, Cochrane Central, Web of Science</p> <p>≤ 2017</p>	<p>patient-reported outcome'/de OR 'patient reported outcome measure'/de OR (('self report'/de) AND ('outcome assessment'/de OR 'quality of life'/exp OR 'quality of life assessment'/exp OR 'complication'/de OR 'symptom'/de OR 'wellbeing'/de OR 'psychological well-being'/de OR 'health status'/de)) OR PROM OR PROMs):ab,ti)</p>	34 (2311)	No HCP criteria specified	Yes

Author Name in Chronological Order	Objectives	Phenomenon of Interest Pertinent to this Review And Clinical Area	Types of Evidence Included and Synthesis Method	Databases Searched and Search Dates	Search Terms for Patient-Reported Measures	Included # Articles (Search Results)	Inclusion Criteria Pertaining to HCPs	Quality or Bias Assessment Conducted
Gelkopf 2020 <sup>1,2</sup>	<p>1. To ascertain the goals of the PROM/ROM projects.</p> <p>2. To identify the setting, diagnoses and services provided in PROM/ROM projects.</p> <p>3. To identify the instruments used to measure outcome,</p> <p>the domains measured and the process that led to the choice of instruments.</p> <p>4. To gather information on the frequency of evaluation and the methods of data collection.</p> <p>5. To gain information about the feasibility of ROM through the follow-up data collection rate; the training needs and type of feedback provided.</p> <p>6. To gather information on the barriers and limitations encountered to implement ROM and ways to overcome those barriers (p. 2).</p>	<p>Barriers and limitations encountered to implement</p> <p>ROM</p> <p>Area: Adult mental health</p>	<p>Peer-reviewed publications including specific initiatives or implementation of PROM data</p> <p>Unnamed, categories with consensus and triangulation</p>	<p>PsyclINFO</p> <p>PubMed (including MEDLINE, Biomed Central, Embase Psychiatry, Elsevier's Science Direct) (cited by Roe, Mazor &amp; Gelkopf, 2019)</p> <p>2000-2018</p>	<p>14 synonyms (cited by Roe, Mazor &amp; Gelkopf, 2019)</p>	<p>103 (19589)</p>	<p>Exclusion of articles on the views, knowledge, attitudes and perceptions of consumers, providers, or other stakeholders regarding the use of PROMs/ROMs.</p>	<p>Not specified</p>

<sup>1</sup>PROMs or PROs

<sup>2</sup>Routine outcome monitoring

The lack of apparent evidence regarding implementation methods and knowledge translation strategies that are most effective across practice areas justifies the need for our proposed systematic review. According to evidence-based practice guidelines for systematic reviews (27, 28) and electronic searching strategies (29), limitations of a systematic review can be traced, for example, to the review's objectives/questions, search strategy, or selection criteria (i.e., inclusion and exclusion). First, getting the question right is of utmost importance in setting up and directing the entire review (27, 30). It was identified that past reviews did not commonly employ the *population, issue/interest, context/comparison, and outcome* (PICO) format and often had several objectives in addition to PCM implementation by HCPs. Inter-related yet divergent objectives may have influenced the search results for relevant citations putting into question the overall validity and thoroughness of the synthesis results (30).

Search terms in combination with relevant databases are fundamental to conducting a literature search at the appropriate level of specificity (i.e., precision and recall). This often elusive balance, when struck, provides the appropriate data to answer the review question (30). Search terms in previous reviews (see Table 1) have been narrow in scope thereby eliciting a low yield of relevant citations. For example, one author included less than five keywords for patient-reported measures (21) while others missed numerous synonyms and appropriate related terms to capture the concept of patient-reported measures (5, 10, 24, 25) or excluded publishing the search terms making the search irreproducible (31). Notably, one review included PREM search terms (24). Furthermore, search terms did not appear to capture the process of implementation in its entirety (10, 11, 23-25) and were limited, in some instances to two words: barriers and facilitators (5, 32). Lastly, some reviews (e.g., (24, 25, 31) had limited reporting of evidence-based electronic search strategies for systematic reviews such as truncation, alternate spelling, and proximity locators (29). When systematic review search strategies do not include search terms at an appropriate level of specificity, the retrieval of relevant citations can be threatened due to low yield. Likewise, robust systematic reviews seeking to balance recall and precision in the retrieval process must also attend carefully to database selection (30).

Selecting the right bibliographic databases to recall a maximum number of relevant sources is another critical aspect of the search process (30). In previous syntheses, the number of relevant citations identified and subsequently screened as eligible have been quite low, which may be associated with the databases used (see Table 1). Four reviews searched three or fewer databases with the most common being MEDLINE/PubMed, CINAHL, and PsycINFO (11, 21, 26, 32). Given the nature of the previous systematic review questions, a broader scope of databases aligned with the population (i.e., all disciplines of HCPs) and the phenomena of interest is warranted to result in a higher yield of relevant citations from which to draw conclusions (30).

PICO elements of review question(s) are utilized to formulate selection criteria (i.e., inclusion and exclusion) to screen the resulting citations for relevance and determination of eligibility (30). Explicit criteria based on the review question(s) determines which citations are included in the reviews and mitigates systematic selection bias; both which are necessary for accurate conclusions to be made (33). Regarding the population element to explore PCM implementation in everyday clinical practice, some limitations were noted in previous reviews (see Table 1). Only three studies specified selection criteria about HCPs (e.g., allied health professionals) (11, 21, 32). Another review included a broad range of stakeholders extending beyond clinicians such as commissioners (e.g., local and national policy makers) and health service managers (24). In terms of context, in half of the published reviews (see Table 1) the selection criteria were limited to only specific practice settings (e.g., palliative care (5); cancer care (10, 23, 26), and adult mental health (25).

Equally important to selection criteria are careful considerations of the types of evidence to include in the review so as to adequately represent the implementation experiences of providers (34). The purpose of mixed methods reviews is to conduct a comprehensive synthesis that includes the different types of evidence necessary to inform clinical practice, policy, or organizational decisions (35). The published systematic reviews (see Table 1) have included a mixture of qualitative and quantitative research-designed studies. However, only Gelkopf et al. (25) stipulated the inclusion of initiatives or projects (e.g., quality improvement and knowledge translation) exploring “real-world” PROM implementation. In recent years, there have been increasing numbers of published quality improvement, organization implementation, and knowledge translation projects exploring PROMs implementation in clinical practice. As such, there is a need to include peer-reviewed evidence about implementation projects and expert opinion because these contribute to a breadth and depth of understanding by capturing everyday experiences of end-users and that might confirm or dispute evidence (35). To date, no single review has captured the voice of end-users, the HCPs, vis-à-vis their experiences of implementing individual-level PCM data at the point of care.

This protocol builds on the work of the aforementioned reviews. Given the cumulative evidence concerning the barriers and facilitators faced by HCPs implementing PCMs in everyday practice, our project will be informed by knowledge translation and implementation science. To this end, our aim is to conduct a comprehensive mixed methods systematic review of a wide range of peer-reviewed literature, including all healthcare settings. The *Population, Phenomenon of Interest, Context* (PIC) format (see Table 2) review question guiding the protocol is “How do healthcare providers implement individual-level patient-centred measurement/assessment tools (and the resultant data) as a routine part of their everyday practice?” This includes the following secondary review questions:

1. What are HCPs’ experiences in applying these tools in clinical practice?
2. How do HCPs interpret and integrate these tools to inform patient care?
3. What are the factors (barriers and facilitators) that influence PCM implementation by HCPs at the point of care?

Table 2. Definitions of the review question elements using the PIC format

PIC Element	Definition
	In this review, patients complete a PROM or PREM in a given practice setting/health service area and then HCPs act on the resultant data in the provision of patient care.
Population (P)	<p><i>Healthcare provider(s)</i></p> <p>Refers to individuals from any health discipline or profession that provide direct health services to patients, clients, and/or families. HCPs are regulated or licenced healthcare professional; however, this may vary by country. HCPs may be referred to as clinicians. Common HCPs include but are not limited to registered nurses, nurse practitioners, physiotherapists, occupational therapists, physicians, social workers, dieticians, psychologists, pharmacists, and midwives.</p>
Phenomena of interest (I)	<p><i>Implement individual-level, patient-centred measurement/assessment tools (and the resultant data)</i></p> <p>In a broad sense, <i>implementation</i> is the process or act of making something active or effective. In the context of PROMs other synonyms include: employ, apply or application, utilize, use, integrate, interpret, draw on, make use of, and act on. The act of implementing PCMs in this review also captures the experiences, views, attitudes habits, practices, and routines of HCPs.</p> <p><i>Patient-centred measurement</i> as an umbrella term refers to standardized assessment tools or questionnaires classified as PROMs or PREMs that every patient is eligible to complete. Thus, the questions are completed by the patient and are about outcomes that matter to them. The results are individual level numerical or textual data that indicate the patient's current state or experience.</p>
Context (C)	<p><i>Routine part of their everyday practice</i></p> <p>Is an expression to describe an action that has been taken as a regular or common aspect of the HCP's role. It is usually in relation to a HCP's direct interaction between them and the patient/client/family. For example, in acute care or hospital settings this term used maybe point-of-care or bedside where patient and providers interact on a regular basis, such as daily, weekly or monthly. The context includes any practice setting or type of health service (e.g., acute care versus community care; private versus publicly funded, and community hospitals versus academic centers/teaching hospitals).</p>

## Methods

The present protocol has been registered within the PROSPERO database (registration number CRD42020182904, November 3, 2020) and is being reported in accordance with the reporting guidance provided in the Preferred Reporting Items for Systematic Reviews and Meta-Analyses Protocols (PRISMA-P) statement (36) [see checklist Additional file 1]. The review will be conducted in accordance with the Joanna Briggs Institute (JBI) Joanna Briggs Institute (JBI) methodology for Mixed Methods Systematic Review (MMSR) (35). To find relevant research evidence, the *population*, *phenomenon of interest*, and *context* (PIC) format has been applied to formulate the review questions, devise a search strategy, and guide study selection criteria. The project will utilize an iterative process undertaken primarily by AW with input from members of the research staff and larger research team based on areas of expertise. A written record will be organized according to the Matrix Method (37).

### Search strategy

The proposed search will be conducted in accordance with the checklist for Peer Review of Electronic Search Strategies (PRESS) Guideline for systematic reviews (29) to achieve a balance of recall and precision. With academic librarian support (DD), the search strategy, including all identified keywords will be adapted for each database. An initial limited search of two databases (MEDLINE and CINAHL) was undertaken to identify articles on the topic. Since controlled vocabulary is unique to each database, keywords (rather than subject headings) have been identified as the most reliable approach for suitable recall. We determined that keyword searching, with the generation of all synonyms, plurals, and alternate spellings (e.g., centred and centered), for the phenomenon of interest (i.e., PROM, PREM, and implementation) [see Additional file 2] produced high yield. Titles, abstracts, and keywords of relevant articles were used to assist in the identification of synonyms for each keyword. Although it is common to include a third keyword to represent the population or context elements of the PIC question, we found that it limited the precision of our results. As such, we decided to exclude a third concept for searching and instead include them as part of our selection criteria. Findings from our preliminary search informed the search for the project.

The evidence to answer our question will be retrieved by searching for the published literature between January 2009 and December 2019 using eight databases with the EBSCO platform that covers the subjects of nursing, allied health, health sciences, psychological literature, physical therapy, occupational health, nutrition, kinesiology, and evidence-based reviews (i.e., MEDLINE, CINAHL, PsycINFO, Web of Science, Embase, SPORTDiscus, Evidence-based Medicine Reviews, and ProQuest (Dissertation and Theses)). The search results will be updated mid-2021 to include literature published since the December 2019 cut-off date of our initial search. In a preliminary search of the literature we noted research about PCMs use at the micro level was first noted around 2009. Initially, we established a 10-year date range. We established a 10-year date range, after a preliminary search of the literature noted that research about PCMs use at the micro level was first published around 2009. Upon further investigation we determined that 71% of the relevant records were published from 2016 onwards. This confirmed our decision to use a 10-year date range to limit the yield of records to those relevant to current clinical practice (e.g., electronic medical records) and synthesize evidence based on the current state of the PCM field. Limiters being used include: (a) scholarly/peer reviewed citations, (b) English language, and (c) date range. The two keywords will be utilized in conjunction with appropriate truncation (e.g., asterisks) to include multiple variants using the same truncated root or stem. In some circumstances the boolean "NEAR" or proximity locators will be used to link terms that may not be adjacent (e.g., barrier\* n4 implement\* and facilitat\* n4 implement\*) (30). All identified search terms will be linked using Boolean operators. The boolean "OR" operator



will be used to link search terms as a union for each concept for the purpose of expanding and broadening a search. The interaction of these concept searches will then be combined with "AND" to narrow the search (30). Search histories [see example Additional file 3] from all databases will be retained.

Our exploration will be supplemented by using other searching strategies to carry out a comprehensive search and counterbalance the limits of keyword database searching (30). This includes: footnote chasing (i.e., scanning the references of "keeper" articles), author searching of those publishing extensively in their field, and backward/forward citation searching of related systematic reviews and other seminal articles (e.g., large studies with numerous publications or those frequently cited) (30). Additionally, a search of authors most frequently publishing in the field will be conducted. The ProQuest database will be used to search for eligible dissertations and theses. Upon completion of our database search, we will search the unpublished literature to lessen publication bias and to retrieve difficult to find literature and information regarding implementation projects. To avoid bias, we will use two approaches. First, a judicious examination of the "grey" literature (e.g., research reports, practice guidelines, and user guides) will be conducted using the Google® search engine in 2020 from a university internet protocol address with the most cited search terms of the PIC concepts identified during our database searches. Second, we will access websites of credible organizations, agencies, and associations that may produce and publish knowledge translation documents supporting PCM implementation (30). These websites will be identified by seeking the opinions of experts in the field. A PRISMA diagram (28) will be created to reflect the search strategy and study selection to reveal how the included studies were determined.

### **Types of Evidence**

The MMSR methodology combines an assortment of evidence to create a breadth and depth of understanding of the review questions posed to inform practice and policy. The inclusion of all available evidence, regardless of type, allows for the degree of agreement or discrepancies between sources of evidence as well as validating or triangulating the findings. Various aspects of a phenomenon of interest can be examined and the available data can contextualize the findings (35). Furthermore, given that implementation at the point-of-care requires a variety of knowledge to inform practice, diverse evidence types will be sought. This review will consider peer-reviewed literature: quantitative, qualitative, and mixed methods studies in addition to reviews (i.e., systematic and literature), organizational implementation projects (e.g., quality improvement, knowledge translation project, implementation of PROMs, program evaluation, or pilot/feasibility project), and expert opinion (e.g., an individual, group or learned body that draws on their practical experience and understanding of the knowledge). We will include not only the evidence on the effectiveness of strategies for implementing PCM ('knowing *what*' type of evidence), but also evidence related to subjective experiences, attitude, behaviours and/or the accepted discourse at the time of practice ('knowing *how*' type of evidence) (34). Opinion-based evidence will be included when derived from expert opinions. That is, the opinions from experts in the field that were gained through some form of consensus building process (e.g., conference, think tank, special interest group, panel, and current discourse) (34). Inclusion of the unpublished grey literature is unique to a systematic review of this nature.

### **Selection Criteria**

The next step to finding relevant evidence for inclusion in this review is to define the selection criteria. To be included in the review, the literature needs to meet the following eligibility criteria (see Table 3): (a) HCPs in a clinical setting; (b) information pertaining to PROM or PREM implementation (e.g., HCP experiences, strategies for integrating into practice, influential factors, or attitudes towards use); (c) data were at the individual level; and (d) any study design. Articles will be excluded if (see Table 3): (a) articles focused exclusively on decision-makers (e.g., managers) or patients; (b) information pertaining to when and why PROMs and PREMs are used as well as impact of their use; (c) studies about instrument development, testing and selection; or (d) implementation of aggregated data. Furthermore, to determine inclusion of studies in the review we will apply the criteria in a specific order (Gough). After each citation is confirmed as written in English and within the date limit, we will ensure it meets the study design criterion. Next, the phenomenon of interest criterion will be applied followed by screening for population and context.

Table 3. Selection Criteria

Topic	Inclusion	Exclusion
Population (P)	<ul style="list-style-type: none"> <li>Healthcare providers</li> </ul>	<ul style="list-style-type: none"> <li>Decision-makers exclusively</li> <li>Patients exclusively</li> </ul>
Phenomena of interest (I)	<ul style="list-style-type: none"> <li>Studies about PREMs or PROMs and               <ul style="list-style-type: none"> <li>experiences of applying or implementing</li> <li>methods or strategies for integrating and interpreting (e.g., processes, logistics, tools, or workflow)</li> <li>factors (barriers and facilitator) influencing implementation</li> <li>views or attitudes towards their use</li> </ul> </li> </ul>	<ul style="list-style-type: none"> <li>Studies about PREMs or PROMs and               <ul style="list-style-type: none"> <li>impact or effectiveness</li> <li>mechanisms by which they work (e.g., patient-provider communication)</li> <li>ways used (e.g., screening, assessment, improve communication)</li> <li>measurement development, testing, and selection</li> <li>suitability for specific patient populations</li> <li>a focus solely on patient-centred care</li> </ul> </li> </ul>
Context (C)	<ul style="list-style-type: none"> <li>Studies concerning data at the individual (micro) level with patients:               <ul style="list-style-type: none"> <li>routine clinical care</li> <li>point-of care</li> <li>everyday clinical practice</li> <li>directly inform patient care or care planning</li> <li>clinical decision-making</li> <li>real-world application</li> </ul> </li> </ul>	<ul style="list-style-type: none"> <li>Studies concerning aggregated data for purposes such as:               <ul style="list-style-type: none"> <li>performance indicators or accreditation</li> <li>value-based medicine</li> <li>quality improvement or quality control</li> <li>resource allocation, service provision, and economic evaluation</li> <li>clinical registries</li> <li>reimbursement and payer issues</li> <li>benchmarking</li> <li>drug development</li> </ul> </li> </ul>
Study Design	<ul style="list-style-type: none"> <li>Published scholarly work including research, pilot or feasibility projects, evidence-based implementation/quality improvement, systematic reviews, literature reviews, and expert opinion</li> </ul>	<ul style="list-style-type: none"> <li>Published literature such as editorials, opinion or position papers, commentary, study protocols, conference proceedings or abstracts, and theory.</li> <li>Insufficient information reported on study design</li> </ul>

### Study Selection Procedure

Following the completion of these searches, all identified citations will be loaded into EndNote X9© (Version 9.3.3) (38) and duplicates removed. AW will provide a thorough orientation to those involved in the selection process to ensure rigor. Given the large quantity of records anticipated to be retrieved (e.g., greater than 20000) the first 100 record titles and abstracts will be screened by two independent reviewers for assessment against the inclusion criteria to be identified as relevant, not relevant, or maybe relevant. Following that, AD will screen all other records to determine relevancy. AW to confirm eligibility will rescreened those identified as potentially relevant. To ensure validity of the selection criteria, in the EndNote© library for this project, AW will conduct keyword searches (e.g., outcome measure, patient outcome, patient-reported) of record titles of those deemed irrelevant to reapply the selection criteria. Finally, all relevant studies will be retrieved in full text and their citation details independently reviewed (AW and AD) against the selection criteria to confirm inclusion. Reasons for further exclusion of all studies will be recorded. Any disagreements that arise between the reviewers at each stage of the study selection process will be resolved through discussion (33, 37).

### Assessment of Methodological Quality

Critical appraisal of included studies will determine the level of evidence and methodological quality as a basis for our confidence to act on the recommendations from our synthesis. Two independent reviewers, blinded to each other's assessments, will retrieve all included citations, and applicable supplemental files, in full text format for assessment. AW will provide a robust orientation to primary reviewers (AD, DG, SH, FH, LE, SL, LM, and two undergraduate research assistants) of the process and appraisal checklists to ensure rigor. Authors will be contacted to request missing or additional data for clarification, where required. Any disagreements that arise between the reviewers will be resolved through consensus discussions among select team members, or a blinded third reviewer.

We will use the following standardized JBI critical appraisal instruments for assessing quality (see Table 4): systematic review, qualitative, cross-sectional, prevalence, case report, and text and opinion (27, 39). To evaluate organizational implementation projects, three questions from the JBI case report checklist (40) were combined with questions from the Johns Hopkins' organizational experience checklist for non-research evidence (41) and questions for quality improvement interventions (42, 43). Similarly, the JBI checklists for analytical cross-sectional (40) and prevalence survey (44, 45) studies were modified to

include four additional questions about the research questions, research methods, ethical approval, and justified conclusions (46-48). JBI checklists do not exist for mixed methods studies or literature reviews. AW conducted an extensive review of the literature to locate other standardized tools of high reliability and validity. Based on a parsimonious set of core criteria, the mixed method checklist focuses on both the effective integration of the quantitative and qualitative components of studies, as well as the provision of a rationale for using a mixed methods design (49-51). The checklist used for literature reviews will be based on the Johns Hopkins' form for non-research evidence (41).

Table 4. Summary of the Critical Appraisal Checklists by Research Design and Level of Evidence

Type of Evidence	Critical Appraisal	Level of Evidence
Systematic review	JBI Systematic Review Appraisal Tool (52)	1
Qualitative	JBI Qualitative Appraisal Tool (53)	3
Analytical Cross-sectional	JBI Analytical Cross Sectional Appraisal Tool with others (above)	3
Survey	JBI Prevalence Appraisal Tool in combination with others (above)	3
Mixed method	Mixed Method Appraisal Tool (51)	3
Organizational implementation	JBI Case Report with others (above)	4
Expert opinion	JBI Text and Opinion Appraisal Tool (39)	5
Literature Review	JH Non-research Evidence (Literature Review) Appraisal Tool (41)	5

All checklists contain a series of criteria (range 8 to 15 questions) scored as being “met” or “not met” or “unclear” and, in some instances, as “not applicable.” Following critical appraisal, all studies will be given a percentage score with higher scores indicating a greater percentage of the quality criteria were met. The research team decided not to set a quality threshold to exclude evidence. Rather, once the data are synthesized we will determine the confidence to act based on the quality and level of the evidence. A modified version of the JBI levels of evidence (54, 55) for meaningfulness will be used as it best aligns with our review questions and the nature of the evidence. The five levels are:

1. Quantitative or mixed-methods systematic review;
2. Qualitative or mixed-methods synthesis and single experimental-based quantitative study;
3. Single qualitative and descriptive or observational quantitative study;
4. Systematic review of expert opinion and organizational implementation project single study (e.g., evidence-based practice, quality improvement, and knowledge translation); and
5. Expert opinion and literature review.

Table 4 cross-references the types of evidence with levels of evidence. Each citation will be assigned a level during the extraction process to be subsequently used during data synthesis. Given the evidence in this review is explorative, descriptive, and interpretative in nature, the JBI Grades of Recommendation will be the criteria used to define the overall strength of the recommendation (i.e., strong or weak) (55, 56).

### Data Extraction

The data extraction step provides the means by which the most pertinent information about the topic (i.e., study characteristics and findings) can be summarized and culled from the primary studies. All source documents will be loaded into the data management software NVivo™ (Version 12.6) (57). Using this software, a Review Matrix will be generated to maximize efficiency and create “order out of chaos” (37, p. 150). Column topics for the matrix will be defined according to the purpose of the proposed systematic review to capture pertinent bibliographic information, methodological characteristics, and content-specific characteristics (e.g., implementation theory) of each included citation (see Table 5) (37). Column topics for which there is a discrete response option (e.g., methodology) will be extracted using the NVivo™ file classification function. The NVivo™ codes function will be used to identify the column topic response for items that have more than one response option (e.g., studies conducted in multiple settings or involving multiple HCPs). These data will offer contextual and methodological data to support the data synthesis results (37). Select team members will be involved in assembling the extracted data from all included articles with relevant accompanying illustrations (e.g., participant quotes or statistical test values). Notes on the definition of column topics and response options as well as the overall extraction process will be kept to ensure consistency amongst extractors.

Table 5. Bibliographic Information and Study Attributes to be Abstracted

Bibliographic Information	Study Attributes
· Authors	· Country/ies of study
· Year of publication	· Methodology
· Article title	· Research design
· Keywords	· Implementation theory
· Digital object identifiers	· Context or Setting
	· Practice area
	· Sample population/profession of healthcare providers
	· Sample size
	· Sampling method
	· Level of evidence
	· PROM and PREM instruments used

The next step will be the extraction of the pertinent study findings, specifically from the results and discussion sections of each citation. Using NVivo™ (57), the process of synthesis begins as the study findings will be extracted into specific codes. All study findings from the included citations will be coded for analysis as textual descriptions. Qualitative data will be composed of themes or subthemes with corresponding illustrations (e.g., quotations, tables, and figures). The quantitative data (e.g., descriptive or inferential statistics) will be converted into “qualitized data.” This process will involve the transformation of all quantitative data into textual descriptions or narrative interpretation in a way that answers the review questions. When necessary, corresponding statistical test results can be captured as part of the coding process. As per the narrative synthesis approach (58), code names will be based on a theoretical framework. In our study, we will use the Consolidated Framework for Implementation Research (CFIR) (59). The CFIR is an evidence-based guideline used to assess multiple contexts and identify factors that might influence the process and effectiveness of the implementation of a specific intervention, which, in our review, is PCM. The five major domains are intervention characteristics (8 items), inner setting (5 items), outer setting (4 items), characteristics of individuals involved (5 items), and implementation process (4 items) (59). A further framework will be used to code the identified implementation processes or actions to support a practice change. For this, we will use the validated Expert Recommendations for Implementing Change (ERIC), which is a compilation of 73 discrete strategies in nine clusters (60, 61). Codes not represented in either framework will be created, as determined, by AW to answer the review questions. In this manner, extraction and initial synthesis occur simultaneously. To reduce coding error during data extraction, we will develop a coding protocol, provide coder training, leverage our substantive expertise amongst team members, and use the NVivo coding comparison feature to improve reliability (30). In summary, the overall extraction process of transforming and coding these data will facilitate each element of the narrative synthesis to integrate the existing evidence and answer the review questions (35).

## Data Synthesis

The synthesis will follow a convergent integrated approach as per the JBI methodology for MMSR. In this manner, data from all types of evidence will be simultaneously extracted and synthesized into meaningful codes. Furthermore, this integrated approach means that the transformed “qualitized” data will be combined to identify patterns across all the studies as well as explore relationships of the data between and within the studies (35). The integration of these data will be guided by a narrative synthesis approach (58), which is well suited for MMSR that utilize diverse types of evidence and has sample heterogeneity (35). Moreover, this approach allows for the use of theoretical frameworks to shape the analysis. In our case, the analysis will use two implementation science frameworks allowing us to focus broadly on the implementation process as well as effective strategies to implement and sustain changes in HCP’s behaviour (58). Popay et al. identifies four iterative elements to a narrative synthesis.

- Element 1: *The role of theory in evidence synthesis* (p. 12). Contributing to knowledge translation theory on how PCM implementation works, why, and for whom we will use the CFIR (59) and ERIC strategies (60, 61) that are based on theories of change. With the use of NVivo™ for extraction, the process of synthesis begins as the theory contributes to the interpretation of study findings and determines how widely applicable the findings may be. In this way theory building and theory testing can be incorporated as a key aspect of the proposed systematic review (58).
- Element 2: *Developing a preliminary synthesis* (p. 13). A preliminary synthesis is conducted to understand the codes identified and summarize the results of included studies. This will be achieved by defining patterns of findings simultaneously across all the studies. An initial description of the findings will evolve based on similarity in meaning to produce an integrated synthesis (58). Furthermore, using NVivo™ to identify codes within each citation will subsequently lead to overarching categories and themes.
- Element 3: *Exploring relationships of the data between and within the studies* (p. 14). The purpose of the third element is to identify reasons that might explain any differences in the findings regarding the successful implementation of PCMs. The emerging patterns identified in the pooled data will be further analysed to identify factors, study characteristics, and context explaining differences. Comparing and contrasting relationships across studies is important to this stage of the synthesis as a means to explore the influence of heterogeneity (58). The NVivo™ relationship and query features will aid in our exploration of associations.
- Element 4: *Assessing the robustness of the synthesis* (p. 15). This element allows for the integration of the quality assessments to determine the strength of the evidence and support with the trustworthiness of the synthesis products (e.g., answers to the study questions and recommendations). Using

NVivo™ the included studies will be assigned both a level of evidence, and a quality score that will be cross-linked to the products of the synthesis. From this, a final determination of the strength of the evidence to support conclusions drawn from the synthesis process can be made (58).

Thus, the results of this narrative synthesis approach will provide a critical analysis to determine effective methods for PCM implementation by HCPs in everyday practice.

## Conclusion

The protocol for this systematic review meets an internationally clinically driven need to conduct a comprehensive synthesis of various types of evidence on (a) effective methods for HCP interpretation and integration of individual-level PCM data, (b) experiences of HCPs applying these tools, and (c) identification of relevant factors influencing PCM implementation. This systematic review is one part of a larger three-phased project that will be followed by HCP interviews and a consensus-building forum to elicit input from key knowledge users. The outcome of the overall project is the development of evidence-informed guidelines that outline effective implementation strategies for integrating PCM data into HCPs' everyday practice for shared decision-making and care planning with patients. These guidelines further build on other user guides for the implementation of PROMs in clinical practice (7, 62, 63). Overall, this project is intended to promote "shared ownership" of individual-level PCM data to better understand and address what matters to patients.

## List Of Abbreviations

CBOs	Clinician-based outcomes
CFIR	Consolidated Framework for Implementation Research
CINAHL	Cumulative Index to Nursing and Allied Health Literature
EBSCO	Elton B. Stephens Company
EMBASE	Excerpta Medica Database
ERIC	Expert Recommendations for Implementing Change
HCP	Healthcare provider
JBI	Joanna Briggs Institute
MEDLINE	Medical Literature Analysis and Retrieval System Online
PCM	Patient-centred Measures
PIC	Population, phenomenon of interest, and context
PICO	Population, issue/interest, context/comparison, and outcome
PREMs	Patient reported experience measures
PRESS	Peer Review of Electronic Search Strategies
PRISMA	Preferred Reporting Items for Systematic Review and Meta-Analyses
PRISMA-P	Preferred Reporting Items for Systematic Review and Meta-Analyses Protocols
PROs	Patient reported outcomes
PROMs	Patient reported outcome measures
PsycINFO	Psychological Index
ROM	Routine outcome monitoring
SPORTDiscus	None available

## Declarations

### Ethical approval and consent to participate

Certificate of approval received from the Trinity Western University Human Research Ethics Board (File No. 19F08)

### Consent for publication

Not applicable

## Availability of data and material

Not applicable

## Competing interests

The authors declare no competing interests.

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## Author's contribution

AW conceived, designed and leads all aspects of the three-phase study, and coordinated the study team. AD assisted with study coordination and participating in protocol steps. SH, DG, LE, and SL assessed methodological quality of included articles. All authors all assisted with the protocol design. All authors read and approved the final manuscript. Each author contributed important intellectual content during manuscript drafting or revision and accepts accountability for the overall work.

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## Authors' information

Development of the review questions were further guided by province-wide stakeholder consultations to generate priority themes for PCM projects. The theme to develop effective knowledge translation strategies for PCM and guidelines on how to interpret and "act on" PCM data was one of the 11 priorities identified.

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## Additional Files

**Additional File 1: PRISMA-P Checklist** (file extension .docx)

Below is a summary of the items addressed in the proposed protocol according to Shamseer et al. (36).



Section and topic	Item No	Item Addressed	Checklist item
<b>ADMINISTRATIVE INFORMATION</b>			
Title:			
Identification	1a	Yes	Identify the report as a protocol of a systematic review
Update	1b	Not applicable	If the protocol is for an update of a previous systematic review, identify as such
Registration	2	Yes	If registered, provide the name of the registry (such as PROSPERO) and registration number
Authors:			
Contact	3a	Yes	Provide name, institutional affiliation, e-mail address of all protocol authors; provide physical mailing address of corresponding author
Contributions	3b	Yes	Describe contributions of protocol authors and identify the guarantor of the review
Amendments	4	Not applicable	If the protocol represents an amendment of a previously completed or published protocol, identify as such and list changes; otherwise, state plan for documenting important protocol amendments
Support:			
Sources	5a	Yes	Indicate sources of financial or other support for the review
Sponsor	5b	Yes	Provide name for the review funder and/or sponsor
Role of sponsor or funder	5c	Yes	Describe roles of funder(s), sponsor(s), and/or institution(s), if any, in developing the protocol
<b>INTRODUCTION</b>			
Rationale	6	Yes	Describe the rationale for the review in the context of what is already known
Objectives	7	Yes	Provide an explicit statement of the question(s) the review will address with reference to participants, interventions, comparators, and outcomes (PICO)
<b>METHODS</b>			
Eligibility criteria	8	Yes	Specify the study characteristics (such as PICO, study design, setting, time frame) and report characteristics (such as years considered, language, publication status) to be used as criteria for eligibility for the review
Information sources	9	Yes	Describe all intended information sources (such as electronic databases, contact with study authors, trial registers or other grey literature sources) with planned dates of coverage
Search strategy	10	Yes	Present draft of search strategy to be used for at least one electronic database, including planned limits, such that it could be repeated
Study records:			
Data management	11a	Yes	Describe the mechanism(s) that will be used to manage records and data throughout the review
Selection process	11b	Yes	State the process that will be used for selecting studies (such as two independent reviewers) through each phase of the review (that is, screening, eligibility and inclusion in meta-analysis)
Data collection process	11c	Yes	Describe planned method of extracting data from reports (such as piloting forms, done independently, in duplicate), any processes for obtaining and confirming data from investigators
Data items	12	Yes	List and define all variables for which data will be sought (such as PICO items, funding sources), any pre-planned data assumptions and simplifications
Outcomes and prioritization	13	Yes	List and define all outcomes for which data will be sought, including prioritization of main and additional outcomes, with rationale
Risk of bias in individual studies	14	Yes	Describe anticipated methods for assessing risk of bias of individual studies, including whether this will be done at the outcome or study level, or both; state how this information will be used in data synthesis
Data synthesis	15a	Not applicable	Describe criteria under which study data will be quantitatively synthesised
	15b	Not applicable	If data are appropriate for quantitative synthesis, describe planned summary measures, methods of handling data and methods of combining data from studies, including any planned exploration of consistency (such as $I^2$ , Kendall's $\tau$ )
	15c	Not applicable	Describe any proposed additional analyses (such as sensitivity or subgroup analyses, meta-regression)
	15d	Yes	If quantitative synthesis is not appropriate, describe the type of summary planned

Section and topic	Item No	Item Addressed	Checklist item
Meta-bias(es)	16	Yes	Specify any planned assessment of meta-bias(es) (such as publication bias across studies, selective reporting within studies)
Confidence in cumulative evidence	17	Yes	Describe how the strength of the body of evidence will be assessed (such as GRADE)

**Additional File 2: Master List of Synonyms for Key Concepts in the Search Strategy** (file extension .docx)

Keyword searching will include search terms from the two key concepts in the PIC-derived review question: patient-centred measurement and implementation. This list contains all generated synonyms, plurals, and alternate spellings.

**Patient-Centred Measurement**

- "family caregiver\* assess\*" OR
- "family caregiver\* index\*" OR
- "family caregiver\* indicator\*" OR
- "family caregiver\* instrument\*" OR
- "family caregiver\* inventor\*" OR
- "family caregiver\* measur\*" OR
- "family caregiver\* outcome\*" OR
- "family caregiver\* questionnaire\*" OR
- "family caregiver\* scale\*" OR
- "family caregiver\* screen\*" OR
- "family caregiver\* survey\*" OR
- "family experience assess\*" OR
- "family experience index\*" OR
- "family experience indicator\*" OR
- "family experience instrument\*" OR
- "family experience inventor\*" OR
- "family experience measur\*" OR
- "family experience outcome\*" OR
- "family experience questionnaire\*" OR
- "family experience scale\*" OR
- "family experience screen\*" OR
- "family experience survey\*" OR
- "family outcome assess\*" OR
- "family outcome index\*" OR
- "family outcome indicator\*" OR
- "family outcome instrument\*" OR
- "family outcome inventor\*" OR
- "family outcome measur\*" OR

"family outcome questionnaire\*" OR  
"family outcome scale\*" OR  
"family outcome screen\*" OR  
"family outcome survey\*" OR  
"family-reported experience questionnaire\*" OR  
"family-reported outcome measur\*" OR  
"family satisfaction assess\*" OR  
"family satisfaction index\*" OR  
"family satisfaction indicator\*" OR  
"family satisfaction instrument\*" OR  
"family satisfaction inventor\*" OR  
"family satisfaction measur\*" OR  
"family satisfaction outcome\*" OR  
"family satisfaction questionnaire\*" OR  
"family satisfaction scale\*" OR  
"family satisfaction survey\*" OR  
"Health care outcome\* assess\*" OR  
"Health care outcome\* index\*" OR  
"Health care outcome\* indicator\*" OR  
"Health care outcome \*instrument\*" OR  
"Health care outcome\* inventor\*" OR  
"Health care outcome\* measur\*" OR  
"Health care outcome\* questionnaire\*" OR  
"Health care outcome\* scale\*" OR  
"Health care outcome\* screen\*" OR  
"Healthcare outcome\* survey\*" OR  
"Healthcare outcome\* assess\*" OR  
"Healthcare outcome\* index\*" OR  
"Healthcare outcome\* indicator\*" OR  
"Healthcare outcome\* instrument\*" OR  
"Healthcare outcome\* inventor\*" OR  
"Healthcare outcome\* measur\*" OR  
"Healthcare outcome\* questionnaire\*" OR  
"Healthcare outcome\* scale\*" OR  
"Healthcare outcome\* screen\*" OR  
"Healthcare outcome\* survey\*" OR

"Health outcome assess\*" OR  
"Health outcome index\*" OR  
"Health outcome indicator\*" OR  
"Health outcome instrument\*" OR  
"Health outcome inventor\*" OR  
"Health outcome measur\*" OR  
"Health outcome questionnaire\*" OR  
"Health outcome scale\*" OR  
"Health outcome screen\*" OR  
"Health outcome survey\*" OR  
"health status assess\*" OR  
"Health status index\*" OR  
"Health status indicator\*" OR  
"health status instrument\*" OR  
"Health status inventor\*" OR  
"Health status measur\*" OR  
"Health status outcome\*" OR  
"Health status questionnaire\*" OR  
"Health status scale\*" OR  
"Health status screen\*" OR  
"Health status survey\*" OR  
"Patient-centred experience" OR  
"Patient-centered experience" OR  
"Patient-centred outcome\*" OR  
"Patient-centered outcome\*" OR  
"Patient experience assess\*" OR  
"Patient experience index\*" OR  
"Patient experience indicator\*" OR  
"Patient experience instrument\*" OR  
"Patient experience measur\*" OR  
"Patient experience outcome\*" OR  
"Patient experience questionnaire\*" OR  
"Patient experience screen\*" OR  
"Patient experience scale\*" OR  
"Patient experience survey\*" OR  
"Patient outcome\* assess\*" OR

"Patient outcome\* index\*" OR  
"Patient outcome\* indicator\*" OR  
"Patient outcome\* instrument\*" OR  
"Patient outcome\* inventor\*" OR  
"Patient outcome\* measur\*" OR  
"Patient outcome\* questionnaire\*" OR  
"Patient outcome\* screen\*" OR  
"Patient outcome\* scale\*" OR  
"Patient outcome\* survey\*" OR  
"Patient-orientated assess\*" OR  
"Patient-orientated index\*" OR  
"Patient-orientated indicator\*" OR  
"Patient-orientated instrument\*" OR  
"Patient-orientated measur\*" OR  
"Patient-orientated questionnaire\*" OR  
"Patient-orientated scale\*" OR  
"Patient-orientated survey\*" OR  
"Patient-oriented assess\*" OR  
"Patient-oriented index\*" OR  
"Patient-oriented indicator\*" OR  
"Patient-oriented instrument\*" OR  
"Patient-oriented inventor\*" OR  
"Patient-oriented measure\*" OR  
"Patient-oriented questionnaire\*" OR  
"Patient-oriented scale\*" OR  
"Patient-oriented screen\*" OR  
"Patient-oriented survey\*" OR  
"patient-reported experience\*" OR  
"patient-reported outcome\*" OR  
"patient-reported satisfaction\*" OR  
"patient satisfaction assess\*" OR  
"patient satisfaction index\*" OR  
"patient satisfaction indicator\*" OR  
"patient satisfaction instrument\*" OR  
"patient satisfaction inventor\*" OR  
"patient satisfaction measur\*" OR

"patient satisfaction outcome\*" OR

"patient satisfaction questionnaire\*" OR

"patient satisfaction scale\*" OR

"patient satisfaction screen\*" OR

"patient satisfaction survey\*" OR

"People-centred assess\*" OR

"People-centred index\*" OR

"People-centred instrument\*" OR

"People-centred measur\*" OR

"People-centred outcome\*" OR

"People-centred screen\*" OR

"People-centred scale\*" OR

"People-centred survey\*" OR

"patient-centred healthcare outcomes" OR

"patient-centred health care outcomes" OR

"patient-centered healthcare outcomes" OR

"patient-centered health care outcomes" OR

"People-centered assess\*" OR

"People-centered index\*" OR

"People-centered indicator\*" OR

"People-centered instrument\*" OR

"People-centered measur\*" OR

"People-centered outcome\*" OR

"People-centered screen\*" OR

"Person-centered outcome\*" OR

"Person-centred outcome\*" OR

"person-reported outcome\*" OR

"person-reported experience\*" OR

"PREM" OR

"PREMs" OR

"PROM" OR

"PROMs" OR

"PROMIS"

## **Implementation**

"Attitudes of health personnel" OR

Barrier\* n4 implement\* OR  
challeng\* n4 implement\* OR  
"Clinical application\*" OR  
"clinical decision making" OR  
"Clinical decision support\*" OR  
"clinical educat\*" OR  
"clinical support\*" OR  
"clinician attitude\*" OR  
"clinician perception\*" OR  
"clinician view\*" OR  
"computer application\*" OR  
"electronic application\*" OR  
Enabler\* OR  
Experience\* OR  
facilitat\* n4 implement\* OR  
"health personnel attitude" OR  
"Implementation science" OR  
Implement\* OR  
integrat\* OR  
Interpret\* OR  
"Knowledge translat\*" OR  
"online application\*" OR  
"Practice guid\*" OR  
"Practise guid\*" OR  
"Practic\* application\*" OR  
"practice educat\*" OR  
"practice support" OR  
"Professional practice" OR  
"Professional development" OR  
"program evaluat\*" OR  
"provider attitude\*" OR  
"provider perception\*" OR  
"provider view\*" OR  
"quality improve\*" OR  
"smartphone application\*" OR  
"stakeholder engagement" OR

"Training project" OR

"Training program" OR

"training clinician\*" OR

"web\* application\*"

**Additional File 3: Example of Search Strategy in Bibliographic Database** (file extension .docx)

Draft of search strategy to be for CINAHL database with year and language limits.

Database name: CINAHL Complete

Date searched: 2019-12-10

Limiters: Human

Language: English

Type of document: peer reviewed, academic journals

Date range: January 1, 2010 to December 31, 2020



Implementation	AND	PCM
<p>"Attitudes of health personnel" OR  Barrier* n4  implement* OR  challeng* n4  implement* OR  "Clinical  application*" OR  "clinical decision  making" OR "Clinical  decision support*" OR  "clinical educat*" OR  "clinical support*" OR  "clinician attitude*" OR  "clinician  perception*" OR  "clinician view*" OR  "computer  application*" OR  "electronic  application*" OR  Enabler* OR  Experience* OR  facilitat* n4  implement* OR  "health personnel  attitude" OR  "Implementation  science" OR  Implement* OR  integrat* OR  Interpret* OR  "Knowledge  translat*" OR "online  application*" OR  "Practice guid*" OR  "Practic*" OR  application*" OR  "practice educat*" OR  "practice  support" OR  "Practise guid*" OR  "Professional  practice" OR  "Professional  development" OR  "program evaluat*" OR  "provider  attitude*" OR  "provider  perception*" OR  "provider view*" OR  "quality improve*" OR  "smartphone  application*" OR  "stakeholder  engagement" OR  "Training project" OR  "Training  program" OR  "training clinician*" OR  "web*" OR  application**</p>		<p>"family caregiver* assess*" OR "family caregiver* index*" OR "family caregiver* indicator*" OR "family caregiver* instrument*" OR "family caregiver* inventor*" OR "family caregiver* measur*" OR "family caregiver* outcome*" OR "family caregiver* questionnaire*" OR "family caregiver* scale*" OR "family caregiver* screen*" OR "family caregiver* survey*" OR "family experience assess*" OR "family experience index*" OR "family experience indicator*" OR "family experience instrument*" OR "family experience inventor*" OR "family experience measur*" OR "family experience outcome*" OR "family experience questionnaire*" OR "family experience scale*" OR "family experience screen*" OR "family experience survey*" OR "family outcome assess*" OR "family outcome index*" OR "family outcome indicator*" OR "family outcome instrument*" OR "family outcome inventor*" OR "family outcome measur*" OR "family outcome questionnaire*" OR "family 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inventor*" OR "Patient-oriented measure*" OR "Patient-oriented questionnaire*" OR "Patient-oriented scale*" OR "Patient-oriented screen*" OR "Patient-oriented survey*" OR "patient-reported experience" OR "patient-reported outcome*" OR "patient-reported satisfaction*" OR "patient satisfaction assess*" OR "patient satisfaction index*" OR "patient satisfaction indicator*" OR "patient satisfaction instrument*" OR "patient satisfaction inventor*" OR "patient satisfaction measur*" OR "patient satisfaction outcome*" OR "patient satisfaction questionnaire*" OR "patient satisfaction scale*" OR "patient satisfaction screen*" OR "patient satisfaction survey*" OR "People-centred assess*" OR "People-centred index*" OR "People-centred instrument*" OR "People-centred measur*" OR "People-centred outcome*" OR "People-centred questionnaire*" OR "People-centred scale*" OR "People-centred survey*" OR "patient-centred healthcare outcomes" OR "patient-centred health care outcomes" OR "patient-centered 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