Journal of Participatory Medicine

Advancing the understanding and practice of participatory medicine among health care professionals and patients
Volume 11 (2019), Issue 1   ISSN 2152-7202   Editors-in-Chief: Susan Woods, MD, MPH; Matthew F Hudson, PhD, MPH

Contents

Review

Participatory Methods to Engage Health Service Users in the Development of Electronic Health Resources: Systematic Review (e11474)
Gaye Moore, Helen Wilding, Kathleen Gray, David Castle. ................................................................. 2

Original Papers

Application of Community-Engaged Research to Inform the Development and Implementation of a Peer-Delivered Mobile Health Intervention for Adults With Serious Mental Illness (e12380)
Karen Fortuna, Paul Barr, Carly Goldstein, Robert Walker, LaPrincess Brewer, Alexandra Zagaria, Stephen Bartels................................. 33

Meaningful Partnerships: Stages of Development of a Patient and Family Advisory Council at a Family Medicine Residency Clinic (e12105)
Jeffrey Schlaudecker, Keesha Goodnow, Anna Goroncy, Reid Hartmann, Saundra Regan, Megan Rich, Adam Butler, Christopher White. . . . 4

2
Review

Participatory Methods to Engage Health Service Users in the Development of Electronic Health Resources: Systematic Review

Gaye Moore1,2*, BN (Hons), MPH, PhD; Helen Wilding1,3*, GradDipInfoMgt; Kathleen Gray4, PhD; David Castle1,5, MBChB, MSc, MD, DLSHTM, GCUT, FRCPsych, FRANZCP

1Mental Health Executive Services, St Vincent's Hospital, Melbourne, Fitzroy, Australia
2Department of Nursing, Faculty of Medicine, Dentistry and Health Sciences, University of Melbourne, Melbourne, Australia
3Library Service, St Vincent's Hospital Melbourne, Fitzroy, Australia
4Health and Biomedical Informatics Centre, University of Melbourne, Melbourne, Australia
5Department of Psychiatry, Faculty of Medicine, Dentistry and Health Sciences, University of Melbourne, Melbourne, Australia
*these authors contributed equally

Corresponding Author:
Gaye Moore, BN (Hons), MPH, PhD
Mental Health Executive Services
St Vincent's Hospital, Melbourne
PO Box 2900
Fitzroy, 3065
Australia
Phone: 61 392311938
Email: gaye.moore@svha.org.au

Abstract

Background: When health service providers (HSP) plan to develop electronic health (eHealth) resources for health service users (HSU), the latter’s involvement is essential. Typically, however, HSP, HSU, and technology developers engaged to produce the resources lack expertise in participatory design methodologies suited to the eHealth context. Furthermore, it can be difficult to identify an established method to use, or determine how to work stepwise through any particular process.

Objective: We sought to summarize the evidence about participatory methods and frameworks used to engage HSU in the development of eHealth resources from the beginning of the design process.

Methods: We searched for studies reporting participatory processes in initial development of eHealth resources from 2006 to 2016 in 9 bibliographic databases: MEDLINE, EMBASE, CINAHL, PsycINFO, Emcare, Cochrane Library, Web of Science, ACM Guide to Computing Literature, and IEEE Xplore. From 15,117 records initially screened on title and abstract for relevance to eHealth and early participatory design, 603 studies were assessed for eligibility on full text. The remaining 90 studies were rated by 2 reviewers using the Mixed Methods Appraisal Tool Version 2011 (Pluye et al; MMAT) and analyzed with respect to health area, purpose, technology type, and country of study. The 30 studies scoring 90% or higher on MMAT were included in a detailed qualitative synthesis.

Results: Of the 90 MMAT-rated studies, the highest reported (1) health areas were cancer and mental disorders, (2) eHealth technologies were websites and mobile apps, (3) targeted populations were youth and women, and (4) countries of study were the United States, the United Kingdom, and the Netherlands. Of the top 30 studies the highest reported participatory frameworks were User-Centered Design, Participatory Action Research Framework, and the Center for eHealth Research and Disease Management (CeHRes) Roadmap, and the highest reported model underpinning development and engagement was Social Cognitive Theory. Of the 30 studies, 4 reported on all the 5 stages of the CeHRes Roadmap.

Conclusions: The top 30 studies yielded 24 participatory frameworks. Many studies referred to using participatory design methods without reference to a framework. The application of a structured framework such as the CeHRes Roadmap and a model such as Social Cognitive Theory creates a foundation for a well-designed eHealth initiative that ensures clarity and enables replication across participatory design projects. The framework and model need to be clearly articulated and address issues that include resource availability, responsiveness to change, and the criteria for good practice. This review creates an information resource for future eHealth developers, to guide the design of their eHealth resource with a framework that can support further evaluation and development.
Apart from operational needs for HSU participation, there are ethical reasons for it. HSP have an ethical responsibility for ensuring that eHealth innovations achieve health outcomes for HSU. HSP are committed to evidence-based practice, in this as in other aspects of their work. Therefore, when they think about developing and deploying new eHealth resources, where do they find what is recognized as good practice in HSU participation? There are so many case studies that it is a near-impossible task to synthesize them all; furthermore, some talk the talk but do not walk the walk of HSU participation, some do not follow any recognized methodology, and some finish early in the life cycle of the eHealth resource.

There are numerous reviews and design guidelines that generalize about theories and methods of HSU participation in eHealth design. They emphasize the importance of the following basic principles:

- appreciation and understanding from the outset, of the range of potential HSU characteristics, goals, needs, values, and perspectives on use [5-7]
- attention to the needs of HSU not just as individual actors but also within their formal and informal care networks [8]
- careful alignment of diverse concerns, attitudes, and perspectives that expert content creators, HSP, and HSU may have [9-12]
- genuine active involvement so that HSU have an opportunity to identify practical problems and design, test, evaluate, and make decisions about technology in a range of environments [13-15]

At the same time, they note that methods of HSU participation in eHealth design need to use human and other project resources judiciously. The themes are as follows:

- ensuring that complex planning and evaluation models are able to be translated and streamlined to develop resources that are practical, feasible, and impactful in real-life settings [16]
- taking a systematic approach to requirements specification to avoid mismatch with the organizational context and to support summative evaluation on a feature-specific level [17]
- applying automation to expedite routine steps to create libraries of typical users and use cases and to manage unforeseen lessons learned for efficiency [18,19]
Nevertheless, key considerations aside, it is difficult for HSP to identify from the literature a recognized, reliable methodological framework for engaging with HSU in the development of eHealth resources. A recent systematic review found that the literature variously encompassed 6 key phases and 17 different methods of participatory design, and it also found that sufficiency of reporting was poor and that no study undertook a robust assessment of efficacy [20]. This makes it difficult for HSP to study the effects of HSU participation in eHealth resources development on reach, adoption, acceptance, and efficacy of the intervention. Relative to other areas of health research, this type of study is immature, without widely endorsed methodological conventions for describing realistic aims for such projects or for determining valid measures of such effects [21].

Therefore, this paper investigates reports of eHealth applications and tools and resource development to determine what methods have been used systematically to ensure full HSU participation. We sought to distill evidence of positive, negative, or other unanticipated effects that have arisen at any stage in the eHealth resource life cycle from various HSU participation methods. Within these participatory approaches, we identified the reported impact from the point of view of HSU and HSP.

The impetus for this study began when the authors sought a strong research framework within which to undertake co-design of an eHealth initiative. The project was based on a print-based and workshop-based psychoeducational intervention called the Optimal Health Program (OHP). The authors wanted to ensure that they chose a rigorous methodological framework for redevelopment of OHP as an eHealth resource. Utilizing proven participatory methods would (1) optimize HSU engagement with the OHP resource that was developed, (2) strengthen the relevance of the resource to intended HSU, and (3) provide a logical foundation for long-term evaluation and improvement of the resource.

**Objectives**

This paper reviews published research reports that include detailed descriptions of participatory methods to engage HSU in eHealth resource development projects. Through synthesizing answers to the following questions, the objective of this paper is to support critical evaluation of this type of methodology and informed selection of appropriate approaches in future research and development projects:

1. What types of eHealth resources have been developed using participatory processes, intended for what types of end users?
2. What frameworks have been used from the very beginning of the design process to ensure participation by the intended end users in the development of eHealth resources?
3. What methods within these frameworks have been most effective in supporting full involvement by intended end users of eHealth resources?
4. What aspects of the participatory methods in these eHealth projects have emerged as being most important to end users?
5. What positive, negative, or other unanticipated effects of participatory methods have the researchers reported at eHealth resource design, development, implementation, or evaluation stages?

**Methods**

**Protocol and Registration**

This systematic review has been carried out in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement [22,23]. Protocol CRD42017053838 was lodged with the PROSPERO international prospective register of systematic reviews.

**Information Sources**

A total of 9 bibliographic databases were searched, including 6 health and biomedical databases and 3 technology databases:

- Ovid MEDLINE(R) Epub Ahead of Print, In-Process & Other Non-Indexed Citations, Ovid MEDLINE(R) Daily and Ovid MEDLINE(R) 1946 to Present (“MEDLINE”)
- EMBASE (Eembase.com ) (“EMBASE”)
- CINAHL Plus with Full Text (EBSCOhost) (“CINAHL”)
- PsycINFO 1806 to January Week 4 2017 (Ovid) (“PsycINFO”)
- Ovid Emcare 1995 to 2016 week 49 (“Emcare”)
Cochrane Library, including Cochrane Database of Systematic Reviews; Database of Abstracts of Reviews of Effect; Cochrane Central Register of Controlled Trials; Cochrane Methodology Register; Health Technology Assessment Database; NHS Economic Evaluation Database; About the Cochrane Collaboration ("Cochrane")
• Web of Science Core Collection ("Web of Science")
• ACM Guide to Computing Literature ("ACM")
• IEEE Xplore Digital Library ("IEEE")

Additional articles were identified from reference lists of key articles and cited by references in Google Scholar.

Search
Search strategies were developed by an experienced medical research librarian (HW) in consultation with the OHP project leader (GM) and an expert eHealth researcher (KG).

In December 2015, scoping searches were developed and run in MEDLINE, EMBASE, CINAHL, PsycINFO, and Cochrane. In April 2016, brief confirmatory searches were run in Google Scholar to consider gaps in the initial strategy and additional search terms or databases that could be included. As a result, search strategies were refined and rerun in the initial health and biomedical databases as well as 3 additional technology databases: Web of Science, ACM, and IEEE. In February 2017, searches were updated to include results to the end of 2016. At this stage, an additional health database, Emcare, was also searched.

Within the health and biomedical databases (MEDLINE, EMBASE, CINAHL, PsycINFO, Emcare, and Cochrane) the search strategies combined the general concepts of user participation AND electronic resources AND program design. These search strategies were not limited to health-related conditions or resources because they yielded a small proportion of nonhealth-related results that could be removed manually. This enabled a very wide range of health conditions, HSU, organizations, and resources to be included in the results.

Within the broader technology databases that are not health specific (Web of Science, ACM, and IEEE), the search strategies were necessarily limited to health-related resources, combining the general concepts of user participation AND electronic resources AND (health OR well-being) AND program design.

A detailed search strategy was developed for MEDLINE using a combination of Medical Subject Headings (MeSH) and text words (Textbox 1). This was then adapted for the other databases, taking into account relevant subject headings and syntax. Search results were limited to publications dated from January 2006 to December 2016 and publications in English language. All database searches were updated in February 2017. Final search strategies for all databases are provided in Multimedia Appendix 1.

Textbox 1. Search strategy for Ovid MEDLINE.

<table>
<thead>
<tr>
<th>Search strategy for Ovid MEDLINE</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ovid MEDLINE(R) Epub Ahead of Print, In-Process &amp; Other Non-Indexed Citations, Ovid MEDLINE(R) Daily and Ovid MEDLINE(R) 1946 to Present</td>
</tr>
<tr>
<td>1. Community-Based Participatory Research/ or consumer participation/ or patient participation/</td>
</tr>
<tr>
<td>2. (codesign* or co-design* or coproduc* or co-produc* or co-creat* or co-creat* or participatory or e-collaboration or usability or focus group*).ti,ab.</td>
</tr>
<tr>
<td>3. ((user* or patient* or consumer* or family or families or carer* or caregiver* or participant* or client* or stakeholder* or peer*) adj2 (centre* or center* or centric or involv* or participat* or partner* or activat* or experience or advisor* or includ* or inclusion or engag* or collaborat* or consult* or empower* or input* or led or focus*)).ti,ab.</td>
</tr>
<tr>
<td>4. 1 or 2 or 3</td>
</tr>
<tr>
<td>5. internet/ or blogging/ or social media/ or audiovisual aids/ or multimedia/ or cell phones/ or text messaging/ or webcasts as topic/ or Telemedicene/or videoconferencing/ or educational technology/ or audiovisual aids/ or motion pictures as topic/ or multimedia/ or exp optical storage devices/ or radio/ or exp tape recording/ or exp television/ or Mobile Applications/ or Software Design/</td>
</tr>
<tr>
<td>6. (internet or web* or online or www* or audiovisual* or audio-visual* or multimedia or multi-media or health or e-health or mobile tech* or mobile phone* or mobile device* or mobile health or mhealth or m-health or cell phone or cellular phone or smartphone or app or apps or blog* or social media or social network* or facebook or podcast* or tracking device* electronic health device* or fitness or elearning or wearable device* or smartwatch* or wearable electronics or telemedicine or tele-medicine or tele-health or tele-health or video* or electronic patient record* or electronic medical record* or electronic health record* or electronic record* or wiki* or portal* or behavioural intervention technology* or health information technology* or software or medical informatic* or health informatic* or digital health).ti.</td>
</tr>
<tr>
<td>7. 5 or 6</td>
</tr>
<tr>
<td>8. Program development/ or planning techniques/ or equipment Design/ or software design/</td>
</tr>
<tr>
<td>9. (develop* or creat* or plan* or build* or implement* or codesign* or co-design*).ti,ab. or design*.ti.</td>
</tr>
<tr>
<td>10. 8 or 9</td>
</tr>
<tr>
<td>11. 4 and 7 and 10</td>
</tr>
<tr>
<td>12. limit 11 to (english language and yr=&quot;2006 -2016&quot;)</td>
</tr>
</tbody>
</table>
Study Selection
The search results were exported from all bibliographic databases to Endnote bibliographic management software. Duplicates were identified and manually removed within Endnote by HW. The records were initially screened within Endnote on title and abstract by HW, excluding results that were clearly irrelevant, that is, not electronic technology, not health or well-being related, not development processes, or not involving end users. All potentially eligible records were exported from Endnote to Covidence, an online platform for managing the systematic review process. Covidence was used to screen records on title and abstract by any 2 of GM, HW, KG, and 1 additional reviewer using predefined inclusion and exclusion criteria as shown in Textboxes 2 and 3. All types of study design were eligible for initial inclusion.

Full-text articles were obtained and uploaded to Covidence for all the available records that had been included based on title and abstract. When a number of articles reported on the same project, they were grouped into a single study to be reviewed together. The full text was reviewed independently by 2 reviewers, GM and HW, using additional inclusion and exclusion criteria (Textboxes 4 and 5).

Textbox 2. Inclusion criteria for screening on title and abstract.
- English language
- Author identified
- Abstract available
- Intended end users are health service users (HSU)
- Intended HSU are involved by proxies: patient associations/advocates/family caregivers
- End users over 14 years of age
- End users are involved in the initial design/development stages

Textbox 3. Exclusion criteria for screening on title and abstract.
- Language other than English
- Author not identified
- Abstract unavailable
- Intended end users are health service providers (HSP) only
- Intended end users are patient advocates, informal caregivers, or family caregivers in their own right
- End users under 14 years of age
- End users are involved only in the later stages of development
- Dissertation
- Duplicate record

Textbox 4. Additional inclusion criteria for full-text review.
- Full text available
- Full conference papers
- Sufficient information on early design/development
- Inclusion of end users on their own behalf
- Development of a specific electronic health (eHealth) resource
- The eHealth resource is designed to support HSU interaction
Textbox 5. Additional exclusion criteria for full-text review.

- Full text unavailable
- Conference abstract only
- Insufficient information on early design stage
- Patient associations/advocates/family caregivers as spokespersons for health service users (HSU)
- No specific electronic health (eHealth) resource actually developed
- Text or video content resources only, with no additional interactivity beyond content consumption
- Hardware only
- Research methodology inappropriate
- Project aim unclear
- Review paper only

Data Collection Process

The included studies were found to use qualitative, quantitative, and mixed methods for HSU participation in eHealth resource development; therefore, the Mixed Methods Appraisal Tool (MMAT) Version 2011 [24] was selected to analyze the rigor of these studies. The MMAT’s 19-assessment criteria were added to the extraction stage of Covidence. Each of the included full-text studies was assessed for methodological quality and rated according to the relevant MMAT criteria.

The detail of MMAT ratings under each criterion was extracted and recorded in an Excel spreadsheet. MMAT scoring metrics were used to calculate a total score for each study in order to develop a hierarchy of evidence for the strength of different methodologies (Multimedia Appendix 2).

The 2 reviewers, GM and HW, worked independently using MMAT to assess the methodological quality of papers and minimize risk of bias in assessing the literature. MMAT ratings and reasoning were compared, and conflicts were resolved through discussions between them.

Risk of Bias in Individual Studies

MMAT scores are typically 100%, 75%, 50%, and 25%. They work on the principle that a mixed-methods study is only as strong as its weakest part. This means that mixed-methods studies that have more criteria to meet (4 qualitative plus 4 quantitative plus 3 mixed method, equaling 11 criteria) could potentially be marked down more easily than studies that are purely qualitative and have fewer criteria to score (4 qualitative criteria only). In order to address this potential bias among study types, a decision was made to include an additional score of 90% to rationalize the difference that occurred between 100% and 75% in mixed-methods studies (Multimedia Appendix 2).

After assessment, studies were grouped by MMAT score and sorted into alphabetical order according to the surname of the first author. Although study numbers were initially used by the reviewers for identification purposes, these have been removed so that there is no confusion about study number and ranking. All studies with the same MMAT score hold equal ranking.

Data Items

The 90 studies assessed according to MMAT are summarized descriptively in a table (Multimedia Appendix 3). First, the data items described in Table 2 were manually extracted from the full text by HW and recorded in Excel for analysis. These results were grouped, tallied, and exported into separate tables according to characteristics of the research scope, such as health area, technology, population, or country of study (Multimedia Appendices 4-7).

Additional descriptive data were extracted from the full text of a subset of included studies, namely 30 studies that scored 90% or higher on MMAT. Data were extracted by HW and GM from the full text of each study using the data items listed in Table 3. These details were grouped, sorted, tallied, and exported into tables that summarize the main methods used to engage HSU in participatory development of eHealth resources.

Methods, frameworks, and processes varied enormously among studies; therefore, a decision was made to allocate all reported methods to the 5 stages of a single framework in order to standardize comparison. The Center for eHealth Research and Disease Management (CeHRes) Roadmap [25] was chosen for this purpose because it was specific to eHealth, highly cited (approximately 400 times between 2011 and 2017), based on the review of many eHealth and development frameworks, process oriented (not just a list of methods but a focus on specific steps), and defined within 5 stages.

Models and theories, participatory frameworks and interventions were extracted from the top 30 studies, and HW subsequently searched for additional mentions of them across the full text of the 90 MMAT–rated studies within Endnote.

Risk of Bias Across Studies

To minimize journal bias, a wide range of bibliographic databases were searched, including those with either a health focus or a technology focus. The search results were limited to English language, which could have created a cultural bias in the studies, although the 90 studies included in the quantitative analysis took place across 21 countries.
Table 2. Data items extracted from 90 studies.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Health area</td>
<td>Main area of health or well-being that the technology addresses; for example, cardiovascular diseases, mental disorders. Defined using MeSH⁶ terms—a controlled vocabulary of hierarchical subject headings from MEDLINE</td>
</tr>
<tr>
<td>Purpose</td>
<td>Purpose of the technology; for example, motivation, self-care, or health education. Defined using MeSH Terms—a controlled vocabulary of hierarchical subject headings from MEDLINE</td>
</tr>
<tr>
<td>Technology type</td>
<td>Identified technology developed; for example mobile app or website. If more than one, all technologies were recorded</td>
</tr>
<tr>
<td>Age group</td>
<td>Age group targeted by the resource (not to be confused by the age group of participants in the development process). Simplified into 3 groups: youth (12-24 years), adult (25-64 years), and aged (65+ years). eHealth⁷ projects aimed at children under 14 years were excluded; therefore, this age group was not included</td>
</tr>
<tr>
<td>Gender specific</td>
<td>Gender specific target of a resource; for example, female only or male only. Not recorded if the resource was inclusive of all genders rather than gender specific</td>
</tr>
<tr>
<td>LGBTQI+ specific</td>
<td>LGBTQI+ specific target of a resource; for example, men who have sex with men. Not recorded if resource was inclusive rather than LGBTQI+ specific</td>
</tr>
<tr>
<td>Cultural/multicultural</td>
<td>Research focusing on a particular culture or across a number of different cultures (for example, Indigenous Australians). Not recorded if culture was not reported as an issue; for example, Swedish research taking place in Sweden with Swedish-speaking participants would not be included unless it was also researched in another country with another language for cross-cultural comparison</td>
</tr>
<tr>
<td>Country where studied</td>
<td>Country where the research took place. If more than one, all are included</td>
</tr>
</tbody>
</table>

⁶MeSH: Medical Subject Headings.
⁷eHealth: electronic health.
LGBTQI+: Lesbian, Gay, Bisexual, Transgender, Queer or Questioning, and Intersex+.

Table 3. Additional data items extracted from the top 30 studies.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specific product</td>
<td>Specific resource developed; for example, named mobile app or website URL</td>
</tr>
<tr>
<td>Models and theory base</td>
<td>Defined structures and models within the project design and delivery, such as Stages of Change</td>
</tr>
<tr>
<td>Participatory frameworks</td>
<td>Defined frameworks involving end users in the development of resources, such as, CeHRes⁸ Roadmap</td>
</tr>
<tr>
<td>Interventions</td>
<td>Specific therapeutic program or guideline, such as Acceptance and Commitment Therapy</td>
</tr>
<tr>
<td>Health service user (HSU) population</td>
<td>Defined end user group for a particular eHealth⁷ project; for example, young people with diabetes</td>
</tr>
<tr>
<td>Health service provider (HSP) population</td>
<td>Defined group of health professionals involved in an eHealth project; for example, mental health clinicians, oncologists</td>
</tr>
<tr>
<td>Teams or groups</td>
<td>Defined teams or groups involved in the development of an eHealth project; for example, leadership team, research group, or advisory group</td>
</tr>
<tr>
<td>Methods</td>
<td>Methods or processes used during the development of an eHealth resource. Includes both participatory and nonparticipatory methods (for example, ethics application and literature search). Participants are identified for some methods (for example, Focus Group [HSU] and Interview [HSP]).</td>
</tr>
<tr>
<td>CeHRes Roadmap stage</td>
<td>Methods sorted into different stages of a defined participatory framework known as the CeHRes Roadmap [25]. The 5 stages include: (1) Contextual Inquiry, (2) Value Specification, (3) Design, (4) Operationalization, and (5) Summative Evaluation.</td>
</tr>
<tr>
<td>Themes/findings (HSU's perspective)</td>
<td>Reported feedback from HSU about the eHealth resource and development process</td>
</tr>
<tr>
<td>Author/researcher recommendations</td>
<td>Reported results, limitations, and recommendations</td>
</tr>
</tbody>
</table>

⁸CeHRes: Center for eHealth Research and Disease Management.
⁷eHealth: electronic health.

The development of eHealth resources is a long process, sometimes taking many years, and many publications only reported a portion of the process, with only a few reporting the entire project up to final evaluation. As conference abstracts and grey literature were excluded in favor of journal articles, sections of the development process may have been reported elsewhere but not included in our evaluation. Reference lists and cited by references in Google Scholar were searched with respect to the top 90 studies to locate connected publications reporting later stages of development, but it is possible that
some publications were either missed or published after our review timeframe.

Results

Database searches retrieved 24,674 records, which were exported to Endnote. Duplicates were removed by HW, leaving 15,117 records. These records were screened for broad relevance on title and abstract by HW and 13,096 records were excluded as clearly irrelevant. The remaining 2021 records were assessed for eligibility on title and abstract using the inclusion and exclusion criteria in Textboxes 2 and 3, and 1391 records were excluded.

The 630 remaining records were combined into 603 studies, some of which involved multiple publications. All 603 studies were assessed for eligibility on full text, and 513 studies were excluded according to the criteria in Textboxes 4 and 5, leaving 90 studies for quantitative analysis. During the screening and full text review process, 12 additional records relating to the 90 studies were identified from reference lists or contact with authors, and those records were combined into the studies. See Figure 1 for the PRISMA flow diagram.

A total of 90 studies were assessed for quality according to MMAT. Results are summarized in Table 4 and detailed results are available in Multimedia Appendix 2. An MMAT score of 100% was awarded to 28 studies and 2 studies scored 90%.

Results From 90 Studies Included in Quantitative Analysis

The 8 data items described in Table 2 were extracted from each of the 90 studies (Multimedia Appendix 3).

The major health focus of each study was grouped into a hierarchy of 18 wider MeSH subject headings, summarized in Multimedia Appendix 4. The top 5 health areas were neoplasms (cancer), mental disorders, nutritional and metabolic diseases (including weight management), virus diseases (including HIV), cardiovascular diseases, and endocrine system diseases (including diabetes).

Nine types of technology were reported in the 90 studies, and these are summarized in Multimedia Appendix 5. Websites (56 studies) and mobile apps (32 studies) were the main eHealth technologies developed. Other types of technology reported were decision tools, handheld computers, kiosk applications, personal health records, serious games, wearable devices, and telemonitoring.

Studies targeting specific populations are summarized in Multimedia Appendix 6. Of the 90 studies, 22 (24%) were youth specific, and 9 (10%) focused on the aged. Of the 90 studies, 11 (12%) reported eHealth projects for women only, and 4 (4%) were for men only. Moreover, 3 studies (3%) had a Lesbian, Gay, Bisexual, Transgender, Queer or Questioning, and Intersex+ focus. Fourteen studies (16%) had either a cultural or multicultural focus, such as a bilingual app for Indigenous Australians or the development of a website in both France and Finland.

The 90 studies took place in 21 countries, summarized in Multimedia Appendix 7. The top 6 countries were United States (33 studies), United Kingdom (15 studies), Netherlands (13 studies), Canada (7 studies), Sweden (6 studies), and Australia (6 studies). Studies also took place in Austria, Belgium, Czech Republic, Greece, Denmark, Finland, France, India, Spain, Ireland, Italy, New Zealand, Norway, Republic of Korea, and Saudi Arabia.

Results From 30 Studies Included in Qualitative Synthesis

The 30 studies scoring 90% or higher on MMAT were recorded in Excel spreadsheets and reviewed in detail. Data items listed in Table 3 were extracted for each study (Multimedia Appendix 8).

The 30 studies are listed in Table 5, along with an indication of the CeHRes Roadmap stages reported. There was often a perceived overlap between stages 1 (contextual inquiry) and 2 (value specification) such as when focus groups may have covered both stages at once. Where this appeared to happen, it was reported in the spreadsheet and included in both stages in Table 5. Where the CeHRes Roadmap was particularly useful was in highlighting stages that were often not reported, such as operationalization or summative evaluation (Table 5). It is possible that some of these studies did address each stage but did not report them in journal articles that were reviewed.

A summary of the 30 highest MMAT–rated studies is represented in Table 6 with details of the product developed, technology used and targeted population. The health area and general purpose of each eHealth project, categorized using Medical Subject Headings (MeSH) is summarized in Multimedia Appendix 9.

The methods were recorded in Excel spreadsheets using the original terminology reported in each study. The details included the number of HSU or HSP involved in each process, the order of each activity as reported, and subprocesses within each method (for example, the type of design activity or workshop activity). These details are included in Multimedia Appendix 8. These detailed methods were then grouped so that they could be summarized using a consistent terminology and then compared. This summary of methods is included for each study in Table 7.

Models and theories referred to in the top 30 studies are shown in Table 8.

Tables 9-13 give an overview of the options used to satisfy each stage of the CeHRes roadmap and the popularity of these methods. Many of the methods reported may demonstrate formative evaluation processes occurring as part of an iterative process. We recommend referring to Multimedia Appendix 8 and the original references for additional information that may be able to identify the practical steps that were implemented.
Figure 1. Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) flow diagram.
Lastly, the models and theories, participatory frameworks and interventions identified in the top 30 studies were searched across the full text of the 90 MMAT-rated studies within Endnote, and the results were ranked in order of prevalence in Multimedia Appendices 10-12.

Twenty three models or theories were identified from the 30 studies scoring 90% or higher on MMAT as playing a role in the development of eHealth resources. The most often reported models and theories were Social Cognitive Theory (n=4, 13%) [144], Theory of Planned Behavior (n=3, 10%) [145], Transtheoretical Model (Prochaska Stages of Change) (n=3, 10%) [146], and the Persuasive Technology Theory/Behavior Model for Persuasive Design (n=3, 10%) [147]. A large variety of other models and theories were referred to, with little overlap between studies. Full results are recorded in Multimedia Appendix 10.

A total of 24 named participatory frameworks or approaches were identified from the 30 studies scoring 90% or higher on MMAT (Multimedia Appendix 11). Only 20 of the 30 top scoring studies referred to a specific framework, with many studies referring more broadly to using participatory design or iterative design methods without reference to a particular named framework. The most often reported participatory frameworks or approaches were User-Centered Design ([UCD], n=5, 17%) [148], PAR framework (n=4, 13%) [149], CeHRes Roadmap (n=3, 10%) [25], Medical Research Council (MRC) Guide to Developing and Evaluating Complex Interventions (n=2, 7%) [150-152], and International Patient Decision Aid Standards Collaboration (n=2, 7%) [153].

Some studies referred to specific interventions that were integral to the function of the eHealth resource that was developed. Key interventions identified in Multimedia Appendix 12 were Cognitive Behavior Therapy (CBT), Behavior Change Techniques, and Mindfulness.
<table>
<thead>
<tr>
<th>Study name and references</th>
<th>MMAT score (%)</th>
<th>Contextual inquiry</th>
<th>Value specification</th>
<th>Design</th>
<th>Operationalization</th>
<th>Summative evaluation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ahtinen, 2013 [26]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>__b</td>
<td>—</td>
</tr>
<tr>
<td>Antypas, 2014 [27]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>✓</td>
</tr>
<tr>
<td>Bengtsson, 2014 [28,29]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Buccieri, 2015 [30]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
</tr>
<tr>
<td>Clayman, 2008 [31]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Cordova, 2015 [32]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Dabbs, 2009 [33]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Das, 2013 [34]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
</tr>
<tr>
<td>Davies, 2015 [35,36]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Fennell, 2016 [39,44]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Fonda, 2010 [40,41]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Goldenberg, 2015 [42,43]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Heckman, 2015 [45]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>✓</td>
</tr>
<tr>
<td>Kelders, 2013 [46]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Lubberding, 2016 [37,38,47]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
</tr>
<tr>
<td>Meyer, 2007 [48]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Miller, 2015 [49]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>✓</td>
</tr>
<tr>
<td>Morrison, 2015 [50]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>✓</td>
</tr>
<tr>
<td>O’Brien, 2016 [51]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Peute, 2015 [52]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Revenas, 2015 [53-55]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Sandlund, 2015 [56]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Schnall, 2016 [57]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Skjoth, 2015 [58]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Stinson, 2014 [59]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>✓</td>
</tr>
<tr>
<td>van Bruinissen, 2014 [60,61]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
</tr>
<tr>
<td>Widman, 2016 [62]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>✓</td>
</tr>
<tr>
<td>Winterling, 2016 [63-66]</td>
<td>100</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Ennis, 2014 [67,69]</td>
<td>90</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
</tr>
<tr>
<td>Fleisher, 2014 [68]</td>
<td>90</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>✓</td>
<td>—</td>
</tr>
</tbody>
</table>

bNo information.
Table 6. Product, technology, and population in top 30 studies.

<table>
<thead>
<tr>
<th>Study and references</th>
<th>Product, technology, and population</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ahtinen 2013 [26]</td>
<td>Living application, a wellness app to support physical activity</td>
</tr>
<tr>
<td>Antypas 2014 [27]</td>
<td>Skibotn Rehabilitation Center resource with personal profile, activity calendar, and SMS reminders</td>
</tr>
<tr>
<td>Bucciari 2015 [30]</td>
<td>Supporting and Assisting Youth (SAY) mobile app for homeless youth</td>
</tr>
<tr>
<td>Clayman 2008 [31]</td>
<td>Cancercarelinks.org—Cancer Care Links for women with breast cancer</td>
</tr>
<tr>
<td>Cordova 2015 [32]</td>
<td>Mobile app version of Storytelling for Empowerment (S4E)—HIV/sexually transmitted infections and drug abuse preventive intervention for primary care</td>
</tr>
<tr>
<td>Dabb 2009 [33]</td>
<td>Pocket Personal Assistant for Tracking Health (Pocket PATH) for lung transplant patients</td>
</tr>
<tr>
<td>Das 2013 [34]</td>
<td>Electronic health portal for weight loss patients undergoing treatment</td>
</tr>
<tr>
<td>Davies 2015 [35,36]</td>
<td>Hep B Story—culturally appropriate bilingual mobile app for Indigenous Australians with hepatitis B</td>
</tr>
<tr>
<td>Fennell 2016 [39,44]</td>
<td>Country Cancer Support website</td>
</tr>
<tr>
<td>Fonda 2010 [40,41]</td>
<td>My Diabetes Data Tracker gadget—prototype personal health app for diabetes management</td>
</tr>
<tr>
<td>Goldberg 2015 [42,43]</td>
<td>HIV prevention app for men who have sex with men (MSM)</td>
</tr>
<tr>
<td>Heckman 2015 [45]</td>
<td>Online skin cancer risk reduction intervention for young adults—UV4.me</td>
</tr>
<tr>
<td>Kelders 2013 [46]</td>
<td>Web-based intervention for prevention of depression, based on self-help book Living to the full</td>
</tr>
<tr>
<td>Lubbering 2016 [37,38,47]</td>
<td>Oncokompas—online self-management application for cancer survivors Oncokompas.nl</td>
</tr>
<tr>
<td>Meyer 2007 [48]</td>
<td>studentdepression.org—student focused website for depression self help</td>
</tr>
<tr>
<td>Miller 2015 [49]</td>
<td>Prostate Cancer Online Guide and Resource for Electronic Survivorship (PROGRESS)—Web-based education program for prostate cancer survivors finishing active treatment</td>
</tr>
<tr>
<td>Morrison 2015 [50]</td>
<td>Living well with Asthma—online resource for self-management of asthma</td>
</tr>
<tr>
<td>O’Brien 2016 [51]</td>
<td>LEAP (Living, Eating, Activity, and Planning in retirement)—Web-based lifestyle intervention in retirement</td>
</tr>
<tr>
<td>Peute 2015 [52]</td>
<td>Website for childhood cancer survivors</td>
</tr>
<tr>
<td>Revenas 2015 [53-55]</td>
<td>tRAppen—Swedish app for self-management of physical activity in rheumatoid arthritis</td>
</tr>
<tr>
<td>Sandlund 2015 [56]</td>
<td>Mobile exercise app to prevent falls in senior citizens</td>
</tr>
<tr>
<td>Schnall 2014 [57]</td>
<td>HIV prevention for high risk men who have sex with men (MSM)</td>
</tr>
<tr>
<td>Skjoth 2015 [58]</td>
<td>Web-based decision aid support for pregnant women to make informed choices about Downs Syndrome screening graviditetsportalen.dk</td>
</tr>
<tr>
<td>Stinson 2014 [59]</td>
<td>iCanCope with Pain—mobile based self-management program for youth with chronic pain</td>
</tr>
<tr>
<td>Van Bruinessen 2014 [60,61]</td>
<td>PatientTIME; 3 products: self-directed online communication tool, corresponding evaluation plan, and implementation plan. Empowering patients to communicate with HSP</td>
</tr>
<tr>
<td>Study and references</td>
<td>Product</td>
</tr>
<tr>
<td>----------------------</td>
<td>--------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Widman 2016 [62]</td>
<td>ProjectHeartforGirls.com—interactive Web program to improve sexual</td>
</tr>
<tr>
<td></td>
<td>communication and reduce HIV/sexually transmitted disease risk in</td>
</tr>
<tr>
<td></td>
<td>adolescent girls</td>
</tr>
<tr>
<td>Winterling 2016 [63-66]</td>
<td>Fex-Can, fertility and sexuality following cancer</td>
</tr>
<tr>
<td>Ennis 2014 [67,69]</td>
<td>myhealthlocker - electronic personal health record for people with severe</td>
</tr>
<tr>
<td></td>
<td>mental illness</td>
</tr>
<tr>
<td>Fleisher 2014 [68]</td>
<td>Web-based decision-making intervention in cancer clinical trials PRE-</td>
</tr>
<tr>
<td></td>
<td>ACT (Preparatory Education About Clinical Trials)</td>
</tr>
</tbody>
</table>

*a* MeSH: Medical Subject Headings.

*b* SMS: short message service text messaging.

*c* USA: United States of America.

*d* MSM: men who have sex with men.

*e* UK: United Kingdom.
Table 7. Participatory frameworks and summary of methods in top 30 studies.

<table>
<thead>
<tr>
<th>Study</th>
<th>Participatory framework</th>
<th>Summary of methods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ahtinen 2013 [26]</td>
<td>Constructive design research, persuasive system design model</td>
<td>Interview (HSU), thematic analysis, observation, think aloud, design activity, focus group (HSU), focus group (HSP), co-design workshop (HSU + HSP), affinity wall, magical gadgets, content creation (HSU), iterative development, prototype, prototype evaluation</td>
</tr>
<tr>
<td>Antypas 2014 [27]</td>
<td>—</td>
<td>Focus group (HSU), thematic analysis, prototype, randomized controlled trial (RCT)</td>
</tr>
<tr>
<td>Bengtsson 2014 [28,29]</td>
<td>—</td>
<td>Ethics approval, HSP/expert identification of need, focus group (HSU), focus group (HSP), thematic analysis, literature search, initial mock-up, interview (HSP), iterative development process</td>
</tr>
<tr>
<td>Buccieri 2015 [30]</td>
<td>Spiral technology action research model, youth centered participatory action study</td>
<td>Advisory group (HSU), literature search, focus group (HSU), design workshop (HSU), design activity—sketching idea, content creation (HSU), prototype, prototype evaluation, promotional campaign</td>
</tr>
<tr>
<td>Clayman 2008 [31]</td>
<td>—</td>
<td>HSP/expert identification of needs, interview (HSU)—longitudinal series, thematic analysis, initial mock-up, iterative development process, prototype, prototype evaluation, questionnaire (HSU)</td>
</tr>
<tr>
<td>Cordova 2015 [32]</td>
<td>Agile software development, community-based participatory research, ecodevelopmental framework</td>
<td>Advisory group, focus group (HSU), interview (HSU), thematic analysis, initial mock-up, iterative development process, prototype</td>
</tr>
<tr>
<td>Dabbs 2009 [33]</td>
<td>User Centered Design</td>
<td>Literature search, survey (HSU), interview (HSU), observation, design workshop (HSU), iterative development process, prototype, prototype evaluation, screen capture, usability testing, testing final version, usability questionnaire, RCT</td>
</tr>
<tr>
<td>Das 2013 [34]</td>
<td>Human centered design, iterative participatory design, collaborative analysis of requirements and design</td>
<td>Ethics approval, HSP/expert identification of needs, interview (HSP), observation, thematic analysis, design workshop (HSU), design workshop (HSP), workshop evaluation, design activity—sketching ideas, interview (HSU), co-design workshop (HSU + HSP), iterative development process, prototype, prototype evaluation, think aloud, screen capture, usability testing, questionnaire (HSU), usability questionnaire, implementation at location used for case study</td>
</tr>
<tr>
<td>Davies 2015 [35,36]</td>
<td>Participatory action research framework</td>
<td>Ethics approval, interview (HSU), thematic analysis, focus group (HSU), initial mock-up (storyboard), iterative development process, prototype, translation back and forth, launch event, evaluation questionnaire</td>
</tr>
<tr>
<td>Fennell 2016 [39,44]</td>
<td>Participatory action research framework</td>
<td>Ethics approval, literature search, review other resources, survey (HSU), interview (HSU), thematic analysis, advisory group (HSU), iterative development process, prototype, prototype evaluation, questionnaire (HSU), promotional campaign, launch event, usage statistics collected (google analytics), feedback form/Web survey/follow up survey</td>
</tr>
<tr>
<td>Fonda 2010 [40,41]</td>
<td>User Centered Design</td>
<td>Focus group (HSU), thematic analysis, focus group (HSP), iterative development process, prototype, prototype evaluation, prototype demonstration</td>
</tr>
<tr>
<td>Goldenberg 2015 [42,43]</td>
<td>—</td>
<td>Ethics approval, focus group (HSU), focus group (HSP), interview (HSP), thematic analysis, prototype, prototype evaluation</td>
</tr>
<tr>
<td>Heckman 2015 [45]</td>
<td>—</td>
<td>Survey (HSU), interview (HSU), think aloud, focus group (HSU), focus group (HSP), iterative development process, content development (HSU), prototype, prototype evaluation, readability/health literacy evaluation by experts, cognitive interviewing (HSU), acceptability testing, usability testing, questionnaire (HSU), thematic analysis, pilot testing, RCT</td>
</tr>
<tr>
<td>Kelders 2013 [46]</td>
<td>CeHRes Roadmap, Human Centered Design</td>
<td>Literature search, focus group (HSP), interview (HSU), thematic analysis, rapid prototyping, initial mock up, iterative development process, prototype, think aloud, usability testing, cognitive walkthrough</td>
</tr>
<tr>
<td>Lubberding 2016 [37,38,47]</td>
<td>—</td>
<td>Ethics approval, interviews (HSU), interviews (HSP), thematic analysis, content creation (HSU), iterative development process, prototype, prototype demonstration, think aloud, usability testing, cognitive walkthrough, implementation plan, feasibility study</td>
</tr>
<tr>
<td>Meyer 2007 [48]</td>
<td>Action Research Framework, Action Research Spiral</td>
<td>Ethics approval, advisory group, interview (HSU), survey (HSU), focus group (HSP), content creation (HSU), initial mock up, prototype, questionnaire (HSU), launch event, usage statistics collected, feedback form/Web survey/follow up survey</td>
</tr>
<tr>
<td>Study</td>
<td>Participatory framework</td>
<td>Summary of methods</td>
</tr>
<tr>
<td>-------</td>
<td>------------------------</td>
<td>-------------------</td>
</tr>
<tr>
<td>Miller 2015 [49]</td>
<td>Iterative Design</td>
<td>Literature search, review other resources, HSP/expert identification of needs, focus group (HSU), interview (HSU), initial mock up, content creation (HSU), iterative development process, prototype, observation, think aloud, readability/health literacy evaluation by experts, usability testing, thematic analysis, RCT</td>
</tr>
<tr>
<td>Morrison 2015 [50]</td>
<td>User Centered Design, Medical Research Council Guide to Developing and Evaluating Complex Interventions</td>
<td>Ethics approval, literature search, HSP/expert identification of needs, focus group (HSU), focus group (HSP), thematic analysis, initial mock-up, interview (HSU), iterative development process, prototype, think aloud, RCT</td>
</tr>
<tr>
<td>O’Brien 2016 [51]</td>
<td>Iterative Design</td>
<td>Ethics approval, literature search, HSP/expert identification of needs, co-design workshop (HSU + HSP), design activity, thematic analysis, content creation (HSU), iterative development process, prototype evaluation, cognitive walkthrough</td>
</tr>
<tr>
<td>Peute 2015 [52]</td>
<td>User Centered Design, Iterative Development Methodology, Website Development Model for the Healthcare Consumer</td>
<td>Literature search, survey (HSU), thematic analysis, co-design workshop (HSU + HSP), initial mockup, iterative development process, prototype, think aloud, screen capture, usability questionnaire, heuristic evaluation (experts)</td>
</tr>
<tr>
<td>Revenas 2015 [53-55]</td>
<td>Participatory Action Research Framework, Experience Based Design, User Centered Design</td>
<td>Ethics approval, survey (HSU), focus group (HSU), thematic analysis, advisory group (HSU), focus group (HSP), co-design workshop (HSU + HSP), iterative development process, prototype, observation</td>
</tr>
<tr>
<td>Sandland 2015 [56]</td>
<td>Form-IT, Participatory and Appreciative Action and Reflection, Soft Systems Thinking</td>
<td>Ethics approval, focus group (HSU), interview (HSU), design workshop (HSU), design activity, iterative development process, prototype, prototype evaluation, observation, questionnaire (HSU), thematic analysis</td>
</tr>
<tr>
<td>Schnall 2014 [57]</td>
<td>Information Systems Research Framework, Iterative Design, User Centered Design</td>
<td>Literature search, focus group (HSU), thematic analysis, design workshop (HSU), design workshop (HSP), design activity — sketching ideas, iterative development process, prototype, usability testing, questionnaire (HSU), usability questionnaire, heuristic evaluation (experts)</td>
</tr>
<tr>
<td>Skjoth 2015 [58]</td>
<td>CeHRes Roadmap, International Patient Decision Aid Standards Collaboration</td>
<td>Literature search, focus group (HSU), focus group (HSP), interview (HSU), interview (HSP), observation, thematic analysis, prototype, prototype evaluation, design reflects clinical pathway</td>
</tr>
<tr>
<td>Stinson 2014 [59]</td>
<td>User Centered Design, Iterative design</td>
<td>Ethics approval, survey (HSU), survey (HSP), thematic analysis, focus group (HSU), focus group (HSP), interview (HSU), prototype, RCT</td>
</tr>
<tr>
<td>Van Bruinessen 2014 [60,61]</td>
<td>Intervention Mapping Framework, User Centered Design, Context Mapping Framework, Stanford Guidelines for Web Credibility</td>
<td>Ethics approval, advisory group (HSU), survey (HSU), focus group (HSU), design activity, thematic analysis, iterative development process, think aloud, heuristic evaluation (experts), implementation plan, RCT</td>
</tr>
<tr>
<td>Widman 2016 [62]</td>
<td>—</td>
<td>Interview (HSU), thematic analysis, literature search, review of other resources, advisory group (HSU), focus group (HSU), content creation (HSU), iterative development process, prototype, think aloud, usability testing, RCT</td>
</tr>
<tr>
<td>Winterling 2016 [63-66]</td>
<td>CeHRes Roadmap</td>
<td>Ethics approval, advisory group (HSU), focus group (HSU), interview (HSU), thematic analysis, iterative development process, prototype, RCT</td>
</tr>
<tr>
<td>Ennis 2014 [67,69]</td>
<td>—</td>
<td>Ethics approval, advisory group (HSU), survey (HSU), focus group (HSU), interview (HSU), thematic analysis, prototype, prototype evaluation, feasibility study, HSU as co-facilitators</td>
</tr>
<tr>
<td>Fleisher 2014 [68]</td>
<td>International Patient Decision Aid Standards Collaboration</td>
<td>Survey (HSU), focus group (HSU), interview (HSU), design activity, content creation (HSU), prototype evaluation, observation, RCT, feedback form/Web survey/follow-up survey</td>
</tr>
</tbody>
</table>

a HSU: health service users.
b HSP: health service providers.
c No information.
d RCT: randomized controlled trial.
<table>
<thead>
<tr>
<th>Study and references</th>
<th>Model/theory base</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bengtsson 2014 [28,29]</td>
<td>Common Sense Model</td>
</tr>
<tr>
<td>Clayman 2008 [31]</td>
<td>—(^a)</td>
</tr>
<tr>
<td>Cordova 2015 [32]</td>
<td>Empowerment Theory</td>
</tr>
<tr>
<td>Dabbs 2009 [33]</td>
<td>—</td>
</tr>
<tr>
<td>Das 2013 [34]</td>
<td>—</td>
</tr>
<tr>
<td>Davies 2015 [35,36]</td>
<td>Paasche-Orlow &amp; Wolf’s Model (causal pathways)</td>
</tr>
<tr>
<td>Fonda 2010 [40,41]</td>
<td>—</td>
</tr>
<tr>
<td>Goldenberg 2015 [42,43]</td>
<td>—</td>
</tr>
<tr>
<td>Heckman 2015 [45]</td>
<td>Integrative Model of Behavior Prediction</td>
</tr>
<tr>
<td>Kelders 2013 [46]</td>
<td>Persuasive Technology Theory, Business modeling</td>
</tr>
<tr>
<td>Lubberding 2016 [37,38,47]</td>
<td>—</td>
</tr>
<tr>
<td>Morrison 2015 [50]</td>
<td>Normalization Process Theory</td>
</tr>
<tr>
<td>Peute 2015 [52]</td>
<td>—</td>
</tr>
<tr>
<td>Revenas 2015 [53-55]</td>
<td>—</td>
</tr>
<tr>
<td>Sandlund 2015 [56]</td>
<td>Appreciative Inquiry</td>
</tr>
<tr>
<td>Schnall 2014 [57]</td>
<td>—</td>
</tr>
<tr>
<td>Skjoth 2015 [58]</td>
<td>—</td>
</tr>
<tr>
<td>Stinson 2014 [59]</td>
<td>Social Learning Theory, Behavioral Activation</td>
</tr>
<tr>
<td>Van Bruinessen 2014 [60,61]</td>
<td>—</td>
</tr>
<tr>
<td>Winterling 2016 [63-66]</td>
<td>—</td>
</tr>
<tr>
<td>Ennis 2014 [67,69]</td>
<td>—</td>
</tr>
</tbody>
</table>

\(^a\)No model or theory base described.
Table 9. Methods used in stage 1 (contextual enquiry) of the CeHRes Roadmap—top 30 studies.

<table>
<thead>
<tr>
<th>Process/method</th>
<th>Total studies (n=30), n (%)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advisory group involved (HSU)</td>
<td>6 (20)</td>
<td>[30,32,48,60,61,63-67,69]</td>
</tr>
<tr>
<td>Literature search</td>
<td>10 (30)</td>
<td>[30,33,39,44,46,49-52,57,58]</td>
</tr>
<tr>
<td>Review other resources</td>
<td>2 (7)</td>
<td>[39,44,49]</td>
</tr>
<tr>
<td>HSP(^b) or expert identified needs</td>
<td>6 (20)</td>
<td>[28,29,31,34,49-51]</td>
</tr>
<tr>
<td>Survey (HSU)</td>
<td>9 (30)</td>
<td>[33,39,44,45,52-55,59-61,67-69]</td>
</tr>
<tr>
<td>Survey (HSP)</td>
<td>1 (3)</td>
<td>[59]</td>
</tr>
<tr>
<td>Focus group (HSU)</td>
<td>13 (43)</td>
<td>[27-30,32,40-43,50,53-58,63-67,69]</td>
</tr>
<tr>
<td>Focus group (HSP)</td>
<td>5 (17)</td>
<td>[28,29,42,43,46,50,58]</td>
</tr>
<tr>
<td>Interview (HSU)</td>
<td>13 (43)</td>
<td>[26,31-33,35-39,44,45,47,48,56,62-67,69]</td>
</tr>
<tr>
<td>Interview (HSP)</td>
<td>3 (10)</td>
<td>[34,37,38,42,43,47]</td>
</tr>
<tr>
<td>Observation</td>
<td>2 (7)</td>
<td>[33,34]</td>
</tr>
<tr>
<td>Think aloud</td>
<td>1 (3)</td>
<td>[45]</td>
</tr>
<tr>
<td>Thematic or data analysis</td>
<td>18 (60)</td>
<td>[26-29,31,32,34-44,47,50,52,57,59,62-67,69]</td>
</tr>
</tbody>
</table>

\(^a\)HSU: health service users.  
\(^b\)HSP: health service providers.

Table 10. Methods used in stage 2 (value specification) of the CeHRes Roadmap—top 30 studies.

<table>
<thead>
<tr>
<th>Process/method</th>
<th>Total studies (n=30), n (%)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advisory group involved (HSU)</td>
<td>6 (20)</td>
<td>[30,32,48,60,61,63-67,69]</td>
</tr>
<tr>
<td>Literature search</td>
<td>2 (7)</td>
<td>[28,29,62]</td>
</tr>
<tr>
<td>Review other resources</td>
<td>1 (3)</td>
<td>[62]</td>
</tr>
<tr>
<td>Survey (HSU)</td>
<td>1 (3)</td>
<td>[48]</td>
</tr>
<tr>
<td>Focus group (HSU)</td>
<td>17 (57)</td>
<td>[27-30,32,35,36,40-43,45,49,50,53-56,58-61,63-66,68]</td>
</tr>
<tr>
<td>Focus group (HSP)</td>
<td>5 (17)</td>
<td>[28,29,42,43,48,50,59]</td>
</tr>
<tr>
<td>Interview (HSU)</td>
<td>10 (33)</td>
<td>[26,31,32,37-39,44,46,47,49,58,62,67,69]</td>
</tr>
<tr>
<td>Interview (HSP)</td>
<td>1 (3)</td>
<td>[58]</td>
</tr>
<tr>
<td>Observation</td>
<td>3 (10)</td>
<td>[26,33,58]</td>
</tr>
<tr>
<td>Design workshop (HSU)</td>
<td>3 (10)</td>
<td>[34,56,57]</td>
</tr>
<tr>
<td>Design workshop (HSP)</td>
<td>1 (3)</td>
<td>[34]</td>
</tr>
<tr>
<td>Co-design workshop (HSU+HSP)</td>
<td>2 (7)</td>
<td>[51,52]</td>
</tr>
<tr>
<td>Workshop evaluation</td>
<td>1 (3)</td>
<td>[34]</td>
</tr>
<tr>
<td>Think aloud</td>
<td>1 (3)</td>
<td>[26]</td>
</tr>
<tr>
<td>Design activity</td>
<td>5 (17)</td>
<td>[26,34,51,56,60,61]</td>
</tr>
<tr>
<td>Thematic or data analysis</td>
<td>17 (57)</td>
<td>[26,27,34,37-44,46,47,50,51,53-55,57-62,67,69]</td>
</tr>
<tr>
<td>Content creation (HSU)</td>
<td>1 (3)</td>
<td>[48]</td>
</tr>
<tr>
<td>Rapid prototyping</td>
<td>1 (3)</td>
<td>[46]</td>
</tr>
<tr>
<td>Initial draft or simple mock up</td>
<td>9 (30)</td>
<td>[28,29,31,32,35,36,46,48-50,52]</td>
</tr>
</tbody>
</table>

\(^a\)HSU: health service users.  
\(^b\)HSP: health service providers.
Table 11. Methods used in Stage 3 (design) of the CeHRes Roadmap—top 30 studies.

<table>
<thead>
<tr>
<th>Process/method</th>
<th>Total studies (n=30), n (%)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advisory group involved (HSU&lt;sup&gt;a&lt;/sup&gt;)</td>
<td>7 (23)</td>
<td>[30,32,39,44,53-55,60-66]</td>
</tr>
<tr>
<td>Focus group or group discussion (HSU)</td>
<td>9 (30)</td>
<td>[26,30,40-43,45,53-55,62-66,68]</td>
</tr>
<tr>
<td>Focus group or group discussion (HSP&lt;sup&gt;b&lt;/sup&gt;)</td>
<td>3 (10)</td>
<td>[40-43,53-55]</td>
</tr>
<tr>
<td>Interview (HSU)</td>
<td>10 (33)</td>
<td>[26,31,32,34,39,44,45,49,50,59,68]</td>
</tr>
<tr>
<td>Interview (HSP)</td>
<td>2 (7)</td>
<td>[28,29,37,38,47]</td>
</tr>
<tr>
<td>Design workshop (HSU only)</td>
<td>3 (10)</td>
<td>[30,33,56]</td>
</tr>
<tr>
<td>Design workshop (HSP only)</td>
<td>1 (3)</td>
<td>[57]</td>
</tr>
<tr>
<td>Co-design workshop (HSU + HSP)</td>
<td>4 (13)</td>
<td>[26,34,51,53-55]</td>
</tr>
<tr>
<td>Design activity; for example, card sorting, sketching, affinity wall</td>
<td>7 (23)</td>
<td>[26,30,34,51,56,57,68]</td>
</tr>
<tr>
<td>Content creation (HSU)</td>
<td>7 (23)</td>
<td>[26,30,37,38,47,49,51,62,68]</td>
</tr>
<tr>
<td>Iterative design process</td>
<td>22 (73)</td>
<td>[26,28,29,31-41,44-47,49-57,60-66]</td>
</tr>
<tr>
<td>Prototype, mockup or storyboard</td>
<td>26 (87)</td>
<td>[26,27,30-50,52-59,62-67,69]</td>
</tr>
<tr>
<td>Translation</td>
<td>1 (3)</td>
<td>[35,36]</td>
</tr>
<tr>
<td>Testing/evaluating prototype</td>
<td>14 (47)</td>
<td>[26,30,31,33,34,39-45,51,56,58,67-69]</td>
</tr>
<tr>
<td>Prototype demonstration</td>
<td>2 (7)</td>
<td>[37,38,40,41,47]</td>
</tr>
<tr>
<td>Observation</td>
<td>5 (17)</td>
<td>[33,49,53-56,68]</td>
</tr>
<tr>
<td>Think aloud</td>
<td>9 (30)</td>
<td>[34,37,38,45-47,49,50,52,60-62]</td>
</tr>
<tr>
<td>Screen capture or recording</td>
<td>3 (10)</td>
<td>[33,34,52]</td>
</tr>
<tr>
<td>Readability or health literacy evaluation</td>
<td>2 (7)</td>
<td>[45,49]</td>
</tr>
<tr>
<td>Usability testing</td>
<td>8 (27)</td>
<td>[33,34,37,38,45-47,49,57,62]</td>
</tr>
<tr>
<td>Cognitive walkthrough</td>
<td>3 (10)</td>
<td>[37,38,46,47,51]</td>
</tr>
<tr>
<td>Survey/questionnaire (HSU)</td>
<td>7 (23)</td>
<td>[31,34,39,44,45,48,56,57]</td>
</tr>
<tr>
<td>Usability questionnaire</td>
<td>3 (10)</td>
<td>[34,52,57]</td>
</tr>
<tr>
<td>Heuristic evaluation</td>
<td>3 (10)</td>
<td>[52,57,60,61]</td>
</tr>
<tr>
<td>Thematic or data analysis</td>
<td>14 (47)</td>
<td>[26,28,29,37,38,42,43,45-47,49-57,59]</td>
</tr>
</tbody>
</table>

<sup>a</sup>HSU: health service users.

<sup>b</sup>HSP: health service providers.

Table 12. Methods used in stage 4 (operationalization) of CeHRes Roadmap—top 30 studies.

<table>
<thead>
<tr>
<th>Process/method</th>
<th>Total studies (n=30), n (%)</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Advisory group involved (HSU&lt;sup&gt;a&lt;/sup&gt;)</td>
<td>4 (13)</td>
<td>[30,39,44,60,61,67,69]</td>
</tr>
<tr>
<td>Design reflects clinical pathway</td>
<td>1 (3)</td>
<td>[58]</td>
</tr>
<tr>
<td>Implementation plan</td>
<td>2 (7)</td>
<td>[37,38,47,60,61]</td>
</tr>
<tr>
<td>Feasibility study</td>
<td>2 (7)</td>
<td>[37,38,47,67,69]</td>
</tr>
<tr>
<td>Promotional campaign</td>
<td>2 (7)</td>
<td>[30,39,44]</td>
</tr>
<tr>
<td>HSU as cofacilitators</td>
<td>1 (3)</td>
<td>[67,69]</td>
</tr>
<tr>
<td>Launch event</td>
<td>3 (10)</td>
<td>[35,36,39,44,48]</td>
</tr>
</tbody>
</table>

<sup>a</sup>HSU: health service users.
Discussion

Overview

In the era of digital health, we have a plethora of literature describing the need for better engagement with HSU to improve health care and health services, and we have access to the technologies to create a broad array of websites and mobile apps, but we lack detailed protocols for designing eHealth resources. This systematic review explored the participatory methods and frameworks used to engage HSU in the development of eHealth resources throughout the design process. UCD was most commonly reported but varied in its application and intention. Participatory methods promoting HSU engagement ranged from brief consultation via a review process to genuine collaboration, which included additional responsibility for the HSU in the actual creation process. Research and development projects that describe a conceptual model (such as Social Cognitive Theory) and a structured framework (such as the CeHRes Roadmap, which includes a diagram/flowchart) lay the foundations for us to gain greater insight into how particular processes lead to efficacious and effective eHealth resources.

Electronic Health Initiatives Developed and the Characteristics of Health Service Users

There have been extensive eHealth initiatives to address the issues of accessibility, engagement, health literacy, data collection, health promotion, early intervention, motivation, and behavioral change. Of the 90 MMAT-scored studies, websites and mobile apps make up the majority of eHealth initiatives presented in this review (Multimedia Appendix 5) with a strong multicultural focus (Multimedia Appendix 6). The end users of these eHealth initiatives were young adults, women, and the elderly (Multimedia Appendix 6) with the focus on cancer and mental health (Multimedia Appendix 4). The app has become an engagement tool used by HSP to make health information and health planning more interactive, interesting, and fun for HSU [30,32,35,36,42,43,56]. Moreover, participatory design is thought to enable young people to be creative and have substantial input into the resource development [30].

Participatory Frameworks

Analyzing the procedural frameworks used in our included studies, we found that no 2 studies reported their processes in the same way. The frameworks governing consumer participation were varied with the most reported being UCD, PAR Framework, CeHRes Roadmap, and MRC Guide to Developing and Evaluating Complex Interventions (Multimedia Appendix 11). The methods implemented to seek the HSU perspectives were also varied with the most reported being focus groups, surveys, interviews, prototype/storyboards, think aloud, and literature search (Tables 9-13). Theories and models that influenced procedures most commonly included cognition, behaviors, processes of change, motivation, and empowerment (Multimedia Appendix 10).

The diversity in eHealth initiatives supports creativity, and to ensure validity and strengthen eHealth research, there is a need to integrate a set of protocols for HSU participation and reporting guidelines [154] available via the Enhancing the QUAlity and Transparency Of health Research Network. This would not constrain methodological innovation and would allow a more effective meta-analysis and comparison of participatory development studies.

Methods Used in the Development of Resources

This review looked for evidence of sound methods for engaging HSU during the development of eHealth apps, tools, and resources. We found relatively few reports that described HSU participation throughout development (ie, from contextual enquiry to summative evaluation, Table 5). Furthermore, many of these reports did not provide adequate details according to mixed-methods appraisal standards. As shown in Multimedia Appendix 2, studies out of 603 full texts reviewed met all of our inclusion criteria and scored 90% or higher according to MMAT. This suggests that research training, funding, and dissemination agencies need to attach far greater importance to reports that describe methods more rigorously.

Others have observed that “The diverse communities working in digital health—including government stakeholders, technologists, clinicians, implementers, network operators, researchers, donors—have lacked a mutually understandable language with which to assess and articulate functionality” [155]. Tables 9-13 illustrates how deeply this lack has affected the production of cohesive research evidence, that is, it is virtually impossible to map the semantic relationships among the methodology elements to inform the discourse about what forms of participatory eHealth design work and why. Many methods are generic to human computer interaction, some take a broad behavioral approach and some include methods of measuring health outcomes in the particular area of health where
the intervention is directed. One possible view is that this illustrates a flourishing of innovation and creativity. Another is that this creates a minefield for research training and peer reviewing and may represent a considerable waste of research resources.

Analyzing the conceptual bases for the methods used in the 30 studies scoring 90% or higher on MMAT, we found much variety with 23 different models or theories reported (Multimedia Appendix 10). The most commonly occurring theories were Social Cognitive Theory, Theory of Planned Behavior, Transtheoretical Model, Persuasive Technology Theory, and Health Behavior Theory. This finding offers a sound basis in evidence for future researchers who wish to follow these precedents. However, we note that research in this area has not been informed by other potentially relevant theories (for example, theories that may account better for healthcare consumers’ economic, emotional, or empowerment motives for engagement) [156].

**Effective Involvement of Health Service Users**

This review looked for evidence about the effectiveness of particular approaches in terms of supporting involvement by HSU. Wintering reported strategies implemented to address engagement with HSU, including 1-person central contact, established expectation of roles, compensation for time, reaching a common agreement, and HSU seen as experts on patient perspective [63-66].

It is also possible to reflect on the richness of the findings generated by particular approaches. As shown in Multimedia Appendix 13, each study reported between 2 and 10 major thematic outputs. Reports with relatively concise outputs were Bengtsson [28,29] using participatory research design and O’Brien [51] describing an array of approaches. The most extensive review was reported by Fleisher [68] using the Ottawa Decision Support Framework and participatory design and Goldenberg [42,43] using 3 types of iterative qualitative research approaches. In assessing effectiveness this way, unknown factors may be in play, such as sophistication of the data collection procedures, analytical expertise of the researchers, editorial constraints on reporting results, and temporal pressures on publication.

**Important Aspects of Participatory Methods for Health Service Users**

There were consistent themes that represented HSU priorities in eHealth initiatives across the selected 30 studies represented in Multimedia Appendix 13. Access to relevant, simple, and clear health information was reported consistently across most of the studies highlighting the importance of this information to make informed decisions in a timely manner. A well-designed eHealth resource that includes a framework supporting HSU involvement can significantly impact health literacy for both HSU and HSP. HSU involvement with the development of an eHealth resource created a collaborative process that required transparency and respect as well as clear mediation processes [53-55].

Being involved in the development of an eHealth resource created the opportunity for HSU to clarify the user perspective and support the relevance of the final product. Despite the variety of websites and apps, HSU reported the need for improved access to information, coordination of care, interactivity with information provided, culturally specific information, patient education, and self-management. HSU also acknowledged the importance of confidentiality and privacy when exchanging personal health information over electronic networks.

**Impact of Participatory Methods Reported by Researchers**

The researchers reported a number of key issues highlighting the importance of participatory methods in creating an eHealth resource that was relevant to HSU. In Multimedia Appendix 13, an outline of the research recommendations was documented for the selected 30 studies. Researchers reported on the importance of utilizing a participatory design, which included an iterative process that increased the responsiveness and relevance of their eHealth initiatives. Having the HSUS perspective from the beginning was important as well as ensuring that the process was genuinely collaborative with all participants respected and acknowledged. Utilizing a health behavior theory in combination with a participatory design was noted to enhance the eHealth resource. The theory base acknowledges the importance of motivation, empowerment, and stages of change in supporting the engagement and utilization of the eHealth resource. It was also noted that the eHealth resource needed to be interesting, engaging, and in some instances include a game-playing element. Creating a more positive approach enabled the HSU to engage with serious and difficult health issues and explore options for improved health. Not only did the eHealth resource need to be interesting but it also importantly needed to be intuitive and simple to navigate.

Heckman [45] reported that their eHealth initiative was guided by intervention development, assessment guidelines for behavioral therapy, and health communication programs with health literacy best-practice. Utilizing a participatory design appeared to improve the relevance of the eHealth resource by addressing issues of culture, gender, age, and sexuality (Multimedia Appendix 6). Goldenberg [42,43] reported personalization along with interactive functionality promoted ownership for HSU. A majority of projects included both HSU and HSP in participatory methods across different developmental stages from contextual inquiry to summative evaluation of the project [28,29,31,34-38,40-43,45-55,58-61,63-67,69]. Evaluation is an integral part of participatory methodology; however, this was reported inconsistently across the 30 studies (Multimedia Appendix 8). The inclusion of a standardized tool to evaluate processes and outcomes from the HSU perspective, as part of a participatory framework, may address the need to bring more objectivity to evaluating various studies.

The demand on time and financial resources to implement a participatory design was noted by some researchers [27,30,42,43,57,68]. Availability of resources was an important consideration throughout the design process, which was often iterative. With the rapid change in technology, there is an increasing demand for HSP to be agile and develop eHealth
resources more quickly but still maintaining an evidence-based, best-practice approach inclusive of HSU participation.

**Limitations**
A limitation of our final dataset is that because of the number of papers retrieved, we decided to limit our analysis to published journal articles and to leave out full papers in conference proceedings. It is possible that there are strong participatory processes that have not been reported in detail, or at all, in the journal literature. Moreover, we did not include studies published in languages other than English and therefore we cannot be certain that our dataset reflects work being done around the world.

As our focus was on the inclusion of HSU from the early development process onward, some studies were included that did not extend to a final evaluation of the product, and it was not always possible to consider the success or otherwise of the final eHealth product. As a part of our inclusion criteria, we required some evidence that a specific eHealth product was ultimately created or likely to be taken to completion.

A limitation of our data analysis is that MMAT is a critical appraisal tool to assess the methodological quality of studies. It does not assess the quality of the writing or the content of the research; therefore, it is possible that we have overlooked papers that may be of high quality in other respects but which we have not rated highly here because of the way their methods sections are presented. For example, under MMAT, a paper will not score highly if it does not discuss the impact of the research or report the limitations of a mixed-methods study. The studies may not have rated highly under MMAT if they used both qualitative and quantitative methods but did not acknowledge that this constituted a mixed-methods study or if only selected aspects were reported. For example, a study that reported HSU participation only at the summative evaluation stage may have involved HSU earlier as well, but this would not register in our search results because we looked for descriptions of methods for HSU participation from initial design stages.

Although categorizing all reported methods in these studies according to the 5 stages of the CeHRes Roadmap [25] was a generally useful way to compare processes across studies, absolute consistency was not achievable because of the wide variety of structuring reports, the differing terminology and naming conventions used for similar methods, and the difficulty in allocating all methods accurately to a particular process stage.

**Conclusions**
Agility of eHealth development is problematic in comparison to nonmedical industries as we seek to ensure safety and quality of care for HSU. It is a challenge for eHealth development to follow rigorous methods within a timeframe that responds to current needs, limited resources, and rapid technological changes. Methodological approaches to developing eHealth resources vary but the importance of engaging HSU in participatory design is consistently emphasized. By synthesizing the existing evidence about strong mixed methods for participatory development of eHealth resources, we anticipate that this systematic review will provide others with clearer guidance to plan more rapid and better-structured work of this kind.

**Acknowledgments**
The authors wish to thank Mark Merolli, Honorary Fellow Health and Biomedical Informatics Center, The University of Melbourne for his expertise and assistance. The authors also wish to thank Belinda Muscat and Zali Annersley, students and honorary St Vincent’s Hospital members for their assistance. This study was supported in part by a University of Melbourne Engagement grant.

**Conflicts of Interest**
None declared.

**Multimedia Appendix 1**
Search strategies for all databases.

[PDF File (Adobe PDF File), 239KB - jopm_v11i1e11474_app1.pdf]

**Multimedia Appendix 2**
Scoring of 90 studies according to the Mixed Methods Appraisal Tool Version 2011.

[PDF File (Adobe PDF File), 471KB - jopm_v11i1e11474_app2.pdf]

**Multimedia Appendix 3**
Descriptive summary of 90 studies.

[PDF File (Adobe PDF File), 320KB - jopm_v11i1e11474_app3.pdf]


118. Williamson H, Griffiths C, Harcourt D. Developing young person’s Face IT: Online psychosocial support for adolescents struggling with conditions or injuries affecting their appearance. Health Psychol Open 2015 Jul;2(2):2055102915619092 [FREE Full text] [doi: 10.1136/bmjopen-2016-012423] [Medline: 28070380]


141. Moore et al. JOURNAL OF PARTICIPATORY MEDICINE


Abbreviations

ACM: Association for Computing Machinery
CeHRes: Center for eHealth Research and Disease Management
CINAHL: Cumulative Index to Nursing and Allied Health Literature
eHealth: electronic health
HSP: health service providers
HSU: health service users
IEEE: Institute of Electrical and Electronics Engineers
MeSH: Medical Subject Headings
MMAT: Mixed Methods Appraisal Tool
MRC: Medical Research Council
OHP: Optimal Health Program
PAR: participatory action research
UCD: User-Centered Design

©Gaye Moore, Helen Wilding, Kathleen Gray, David Castle. Originally published in Journal of Participatory Medicine (http://jopm.jmir.org), 22.02.2019. This is an open-access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any
medium, provided the original work, first published in Journal of Participatory Medicine, is properly cited. The complete bibliographic information, a link to the original publication on http://jopm.jmir.org, as well as this copyright and license information must be included.
Application of Community-Engaged Research to Inform the Development and Implementation of a Peer-Delivered Mobile Health Intervention for Adults With Serious Mental Illness

Karen Fortuna¹, PhD; Paul Barr¹, PhD; Carly Goldstein², PhD; Robert Walker³, MS, COAPS; LaPrincess Brewer⁴, MD; Alexandra Zagaria⁵, BA; Stephen Bartels⁶, MD, MS

¹Department of Psychiatry, Dartmouth College, Lebanon, NH, United States
²Brown University, Providence, RI, United States
³Massachusetts Department of Mental Health, Boston, MA, United States
⁴Department of Cardiovascular Medicine, Mayo Clinic College of Medicine, Rochester, MN, United States
⁵Dartmouth-Hitchcock, Lebanon, NH, United States
⁶The Mongan Institute, Massachusetts General Hospital, Boston, MA, United States

Corresponding Author:
Karen Fortuna, PhD
Department of Psychiatry
Dartmouth College
46 Centerra Pkwy, Suite 200
Lebanon, NH, 03766
United States
Phone: 1 6036533430
Email: klfortuna@gmail.com

Abstract

Background: Involving certified peer specialists in all phases of intervention development and research is a high priority to advance peer-delivered services. Certified peer specialists are individuals with a lived experience of a mental illness, and they are trained and accredited to provide Medicaid reimbursable mental health services. Community-engaged research can facilitate the development and implementation of peer-delivered interventions; however, little is known about the processes. We present our application of community-engaged research to inform the development and implementation of a peer-delivered mobile health (mHealth) intervention for adults with serious mental illness.

Objective: The aim of this study was to present a framework that can be used as a guide for researchers and certified peer specialists to develop and implement peer-delivered mHealth interventions in community settings.

Methods: Informed by principles of community-engaged research, we developed the Academic Researchers-Certified Peer Specialists mHealth Research Continuum. Principles of community-engaged research included in the Continuum include the following: (1) develop a clear understanding of the purpose, goal, and population involved in community change; (2) become knowledgeable about all aspects of the community; (3) interact and establish relationships with the community; (4) encourage community self-determination; (5) partner with the community; (6) respect community diversity and culture; (7) activate community assets and develop capacity; (8) maintain flexibility; and (9) commit to long-term collaboration.

Results: Overall, 4 certified peer specialists participated in all phases of intervention development and research. Individuals who participated in the Academic Researchers-Certified Peer Specialists’ mHealth Research Continuum collaborated on 5 studies advancing peers’ roles in services delivery using mHealth and secured grant funding from a foundation to sustain their study. The Academic Researchers-Certified Peer Specialists’ mHealth Research Continuum has created a rare environment of inclusion by combining scientific expertise and certified peer specialists’ expertise to achieve a shared vision.

Conclusions: This study delineates a process by which academic researchers and certified peer specialists participated in community-engaged research to develop and implement peer-delivered mHealth interventions in community settings.

doi:10.2196/12380
KEYWORDS
stakeholder participation; mental health; patient participation; consumer advocacy; mobile health

Introduction

Background

Consumers diagnosed with a serious mental illness (SMI) have been long-established advocates for transparency and full partnerships with providers in treatment settings [1]. These partnerships aim to ensure human dignity, self-determination, and civil rights of consumers with SMI [1]. Despite advocacy efforts, a recent systematic review of peer-delivered intervention studies suggests that certified peer specialists do not significantly interact or assist in intervention development and implementation in a role beyond the interventionists [2]. To our knowledge, this is the first report of a community-engaged research framework that includes certified peer specialists in all stages of research and intervention development. Peers or certified peer specialists are people with a lived experience of a mental illness and have been accredited by the state to provide mental health services such as peer support [3]. Certified peer specialists are part of a national network that offers Medicaid reimbursable peer services in 34 states [4]. Consistent with the national research agenda to advance peer-delivered services [4], we present a community-engaged research framework that includes certified peer specialists in all research stages [4]. This framework can be used as a guide to participate in community-engaged research to develop and implement peer-delivered mobile health (mHealth) interventions in community settings.

Community engagement is defined as “the process of working collaboratively with and through groups of people affiliated by geographic proximity, special interest, or similar situations to address issues affecting the well-being of those people” [5]. A systematic review of community engagement found that most studies that include community-engaged research have a positive impact on health behaviors, such as diet and exercise, as well as health outcomes (eg, obesity, mental well-being, and quality of life). Principles of community engagement set forth by the Clinical and Translational Science Awards Consortium include the following: (1) develop a clear understanding of the purpose, goal, and population involved in community change; (2) become knowledgeable about all aspects of the community; (3) interact and establish relationships with the community; (4) encourage community self-determination; (5) partner with the community; (6) respect community diversity and culture; (7) activate community assets and develop capacity; (8) maintain flexibility; and (9) commit to long-term collaboration [5].

In developing peer-delivered interventions, certified peer specialists represent both interventionists' and the populations' interests. Certified peer specialists, unlike most researchers, have unique expertise and insight into the mental health care system as they have a lived experience utilizing this system for their personal health care needs [6]. Certified peer specialists have the potential to be instrumental members of research teams during all phases of developing behavioral interventions—from idea conception to effectiveness testing. For example, certified peer specialists voice ideas, concerns, and priorities that may not be part of researcher-driven intervention development and implementation. Peers can give guidance on intervention development and research procedures that are acceptable to certified peer specialists, consumers with SMI, and organizations. For example, academic researchers are commonly interested in medical outcomes; however, consumers with SMI and certified peer specialists have expressed an interest in personal recovery outcomes such as hope and empowerment. In addition, incorporating certified peer specialists as equal partners in intervention development and implementation can potentially enhance intervention success [7]. For example, social influences on health are complex and go beyond biologic and health care system factors and include social networks and support systems as well as physical environments [8]. As such, including certified peer specialists can offer valuable perspectives and insights into addressing the needs of similar populations within the context of their social and physical environments.

Objective

Informed by community-engaged research principles, we delineate a process by which certified peer specialists were included as full partners in the development of a peer-delivered mHealth for adults with SMI. The goal of this report is to present a framework that can be used as a guide for researchers and certified peer specialists to develop and implement peer-delivered mHealth interventions in community settings.

Methods

Preliminary Research Before Active Engagement With the Peer Community

Defining the Problem From the Peers’ Perspective

Aligned with the first principle of community-engaged research, academic researchers developed a clear understanding of the social problem experienced by consumers with SMI that certified peer specialists wanted to address [5]. To begin, academic researchers conducted a community assessment using a national survey with 267 certified peer specialists from 38 states. The survey was designed to identify the top 3 biological, psychological, social, or environmental issues confronting people with SMI (Fortuna et al, unpublished data). We used an online survey to engage certified peer specialists with diverse socioeconomic backgrounds to better understand the community’s collective needs. This survey identified the management of mental health and chronic physical health conditions as a major, unaddressed issue (Fortuna et al, unpublished data).

Aligned with the principle of self-determination in community engagement, academic researchers worked with the community’s goal [5] to address mental health and chronic physical health conditions among consumers with SMI. Self-determination theory suggests that individuals have the choice and the right to determine their future [5]. Within community-engaged
research, academic researchers do not have the right to impart academic research needs on the community. The concept of self-determination is consistent with the National and State Peer Support Code of Ethics [9]. Self-determination within community-engaged research is the impetus for community partners to engage in community research [5]. For example, if community members recognize and see value in addressing the problem identified and if they feel they have an influence in decision-making and can make an impact, there is a greater likelihood they will engage. In developing the partnership, academic researchers identified addressing mental health and chronic physical health conditions among people with SMI as the primary goal of the partnership. As such, over the next year, our partnership developed and tested an mHealth intervention designed to address mental and physical health self-management skill development.

**Academic Researcher Capacity Building**

Aligned with the principle of community engagement to become knowledgeable about the community [5], academic researchers developed knowledge of certified peer specialists. Academic researchers’ capacity building began by learning about peers’ beliefs, values, and culture through understanding the history of the mental health care system, reading literature written by peers (eg, *Reaching Across: Mental Health Clients Helping Each Other* [10]), and understanding the Medicaid reimbursement system for certified peer specialists. Although this process was important during early phase knowledge building, working one-on-one with certified peer specialists was the most valuable learning process.

**Mapping Certified Peer Specialists’ Assets**

Next, researchers identified certified peer specialists’ assets. For example, certification as a peer specialist in Massachusetts requires active participation in treatment; completion of an 80-hour training including classes, small group activities, and homework on fundamentals of peer support; cross-cultural partnering; use of first-person, nonclinical language; and passing score on a written examination. Certified peer specialists are then accredited to provide Medicaid reimbursable services in 34 states [4]—most commonly peer support or wellness. As such, there is potential for national dissemination and uptake if the intervention is successful.

In addition, certified peer specialists are a trained workforce with professional practice standards that could guide intervention development. Professional practice standards include (1) not forcing people to participate in services, (2) sharing stories of recovery, (3) not judging others, (4) embracing diversity, (5) educating and advocating for others, (6) addressing difficult issues, (7) learning from people they support and those supported learn from them, (8) embracing equality, (9) using a strengths-based approach, (10) setting clear expectations, and (11) focusing on the person and encouraging them to achieve what they want in life [9].

Finally, nearly all certified peer specialists in the national online survey owned a smartphone (94.8%; 253/267), and everyone indicated that smartphones and tablets could enhance the services they deliver [11]. They reported being willing to deliver smartphone interventions for mental and physical health self-management, suggesting that smartphones may be a useful tool for offering evidence-based care (see Figure 1 for a community map of certified peer specialists’ assets).

**Potential Power Differential**

Next, academic researchers had the opportunity to informally speak with a certified peer specialist who provided services in the Massachusetts area. The purpose of this conversation was to learn more about certified peer specialists. This in-person conversation lasted approximately 30 min. As a result of this informal discussion, academic researchers advanced their knowledge of certified peer specialists beyond written material of peers’ beliefs, values, culture, and history and identified potential power differentials that may impact the relationship. For example, academic researchers learned that peers may be skeptical of people involved in the mental health care system and research—potentially due to historical, structural oppression, and stigma [12]. As such, academic researchers were led by classic and contemporary literature on social justice to assuage this potential power differential.

As academic researchers moved toward formalizing the partnership (see next section: *Early Phase Research Aimed to Foster Certified Peer Specialist Engagement*), principles of fairness, empowerment, inclusion, and self-determination [12-15] were introduced at the forefront of discussions between academic researchers and certified peer specialists. Specifically, academic researchers defined these principles in accordance with well-established definitions [12-15] with certified peer specialists. Then, in an open discussion, academic researchers and certified peer specialists elaborated on these principles and offered examples of how each principle would apply to the partnership. In Textbox 1, we present how the principles of fairness, empowerment, inclusion, and self-determination were incorporated into partnership. The inclusion of these principles formalized a set of guidelines for how the partnership would operate.

**Early Phase Research Aimed to Foster Certified Peer Specialist Engagement**

**Establishing Relationships and Developing Trust**

Aligned with the principle of community-engaged research to establish an authentic relationship with the certified peer specialists’ community [5], academic researchers established a relationship and developed trust. To establish relationships with peer leaders, we developed the first iteration of the mHealth intervention with peers as consultants. The academic researchers contributed scientific knowledge to the design and development of integrated medical and psychiatric self-management interventions and expertise with research methodologies. The academic researchers identified integrated illness management and recovery (I-IMR) as having clinical effectiveness [16] and the possibility of being delivered by certified peer specialists.
Textbox 1. Inclusion of principles of fairness, empowerment, inclusion, and self-determination in mobile health development and implementation.

**Fairness**
Resource sharing: financial incentives, offering transportation assistance, and food; and resource allocation: equitable pay for certified peer specialist

**Empowerment**
Peer training in the research capacity building and de-emphasized that researchers were the experts; rather, both groups brought their unique expertise to the team

**Inclusion**
Full inclusion on research teams and equal credit for mobile health intervention development as evidence by peer-reviewed publications and national presentations with peers as authors or co-presenters

**Self-determination**
Academic researchers worked with the community’s goal and modified the intervention to include peer support in addition to medical and psychiatric self-management skills training and refocused the intervention from the medical model to the recovery model of services delivery

Certified peer specialists did not assist in selecting this intervention to adapt. I-IMR is an evidence-based approach consisting of an individually tailored program addressing physical and mental health self-management in adults with SMI aged over 50 years. The academic researchers selected an intervention for older adults with SMI, as older adults with SMI are more likely to have multiple comorbidities [17] and, thus, a higher need for self-management of both medical and psychiatric conditions. Key to I-IMR is its delivery by a masters-level provider and a nurse over an 8- to 10-month period. There is evidence demonstrating that I-IMR results in improved self-management and decreased hospitalizations in older adults with SMI [16].

The academic researchers adapted I-IMR for in-person delivery by a certified peer specialist. To support fidelity, we included the use of guided eModules (ie, guided curriculum) and a smartphone app designed to complement in-person eModule sessions.

**eModules**
The eModules were designed to be reviewed on a tablet side-by-side with a certified peer specialist and a consumer with SMI during weekly 1-hour in-person sessions in a community setting. Each eModule includes videos and experiential learning tasks on psychoeducation and coping skills training. Academic researchers worked with certified peer specialists and filmed peer-led self-management videos on personal recovery stories (unscripted) that were included in the eModules.

http://jopm.jmir.org/2019/1/e12380/
sessions include (1) Identifying Your Individual Recovery and Wellness Goals: Setting recovery and health goals and strategies to achieve goals and orientation to the smartphone app; (2) Psychoeducation: Psychoeducation on SMI and medical illness; (3) Stress Vulnerability and Illness: Causes of mental illness and factors that influence its course; (4) Building Social Supports and Recovery and Wellness: How to build social supports; (5) Medication Adherence Strategies: Behavioral tailoring and motivational techniques for medication adherence; (6) Psychiatric and Medical Relapse Prevention: Identify warning signs and develop a relapse prevention plan; (7) Coping with Psychiatric Symptoms and Health-related Stress and Solving Problems: Establish a method managing symptoms; (8) Coping with Stress, Chronic Pain, and Medical Symptoms: Identifying stressors that exacerbate symptoms and strategies to cope with stress; (9) Medication Misuse: Addressing medication misuse and the effects on symptoms and functioning; and (10) A Guide to Navigating the Mental Health and Medical Health care System: Accessing mental health and medical health services and making informed decisions.

Smartphone App

The smartphone app was designed to assist in the transfer of self-management skills from in-person sessions with certified peer specialists to real-world environments. As we were developing the smartphone app, we had an informal relationship with 2 certified peer specialists and 2 consumers with SMI. We consulted with these individuals, but they were not part of the scientific team meetings. Our process included asking a clinical social worker to talk to certified peer specialists and consumers with SMI to get their opinions on the features of the smartphone app, but not the content of the intervention. The scientific team consisted of clinical social workers, physicians, and engineers [18]. The app includes personalized (1) homework from in-person meetings, (2) a relapse prevention plan, (3) daily self-management to-do checklist, (4) videos and animations to guide individuals in practicing self-management skills, and (5) articles on self-management. Branching algorithms built into the app technology allow personalization of these features to meet an individual’s personalized recovery goals. The smartphone app also included a chat feature to allow text messaging between certified peer specialists and consumers.

Once we developed these products (app and eModules), we used a scientific approach to allow peers and consumers to evaluate our study and be involved in early technology development. Academic researchers used an adaptive systems engineering approach [19] to conduct a usability test and task analysis (ie, if consumers could use the technology on their own) [18]. The usability test and task analysis was a formal scientific study, in which peers were prompted to provide continuous verbal feedback while using the app and eModules [18]. Peers were free to report whatever they felt relevant. Peers were asked to provide their verbal reactions as they completed tasks on the smartphone (app) and the tablet (eModules). Peers were asked to complete the following tasks: (1) turning the phone and tablet on, (2) finding the icon to launch the program, (3) selecting treatment programs and progressing through each program of the intervention, (4) watching videos, (5) writing and sending a text message using the text message feature within the Web app, (6) finding and checking off daily tasks, (7) reading text on the instruction page, (8) responding to a push notification prompting the participant to watch a video, and (9) returning to the homepage. Peers were asked to engage in all components of the app and eModules and provide feedback on the content, language, layout, colors, typeface, videos, graphics, text size, readability, and navigation features.

During this process, certified peer specialists provided their expertise on the needs of the community and contributed to the context of unhealthy behaviors and poor management of mental and physical health conditions. For example, academic researchers learned that healthy eating is a challenge for individuals with SMI as they may face difficulties with affording healthy foods on a limited income or due to unemployment or underemployment, or they may lack cooking skills or skills to determine healthy eating, lack a reliable place to cook, or they may eat junk food as a means to feel better in the short term. As part of this process, certified peer specialists also identified barriers and facilitators to mHealth implementation and made recommendations to ease intervention uptake. For example, certified peer specialists informed academic researchers that the app should not only be available on a smartphone but also on a tablet because older adults with SMI who wear glasses may experience difficulty reading the small text on the smartphone [18].

At this time, we learned of the need to consider certified peer specialists’ preferences and philosophy of services delivery, social and environmental contexts, and perspectives of feasibility and acceptability from certified peer specialists and consumers with SMI. We needed certified peer specialists to have a more substantive and egalitarian role in the technology development process; we needed to establish a formal partnership and to include certified peer specialists and consumers with SMI in the expert team meetings to move this program of research forward.

Full Academic Researchers-Certified Peer Specialists Partnership

Establishment of a Formal Academic Researchers-Certified Peer Specialist Partnership

Aligned with community-engaged research principles, academic researchers partnered with the certified peer specialist community [5]. We engaged a community gatekeeper to develop a certified peer specialist team to guide the next phases of mHealth intervention development. Community gatekeepers are influential members of the community of interest and provide access to the community [20]. We were introduced to the Consumer Engagement Liaison for the Department of Mental Health in the state of Massachusetts. As the Consumer Engagement Liaison (ie, community gatekeeper) understood the needs of the local community, the Liaison advocated for academic researcher inclusion in the certified peer specialists’ network within the public mental health service system. This community gatekeeper model allowed ease of access and the opportunity to develop a trusting relationship between academic researchers and certified peer specialists.
With the assistance of the Consumer Engagement Liaison, we convened a meeting with certified peer specialists, social workers, disability rights advocates, health services researchers, primary care providers, and the Massachusetts Department of Mental Health. We discussed consumers’ needs with SMI. Together, we established a need for an effective, easily accessible self-management intervention for adults with SMI. Academic researchers presented the adapted version of I-IMR, including the eModules and the smartphone app, newly renamed as PeerTECH. After this initial meeting, a formal research partnership developed with the joint goal of assessing PeerTECH’s feasibility, acceptability, and effectiveness. Overall, 4 certified peer specialists, 2 social workers, 1 disability rights advocate, 3 health services researchers, 2 primary care providers, and 1 individual with the Massachusetts Department of Mental Health participated in all phases of intervention development and research.

Developing an Infrastructure for Full Participation and Shared Decision-Making Authority

Throughout this process, academic researchers respected certified peer specialists’ diversity and culture [5]. We held a series of meetings throughout the pilot study to create an infrastructure that would encourage openness and change (ie, 3 in-person and 13 virtual weekly 1-hour meetings over a 20-week period). We structured the academic researchers-certified peer specialists’ mHealth research group into 2 teams: the scientific team and the peer direct service team. The scientific team met weekly to discuss coordinating the pilot study. The scientific team included 2 people who represented organizational leadership within the selected research site, the community gatekeeper who represented certified peer specialists and consumer interests, 3 social workers familiar with consumer research participants, a peer supervisor (who was also a certified peer specialist), and 2 academic researchers (see Multimedia Appendix 1).

Certified Peer Specialists’ Research Capacity Building

Aligned with the principle of community-engaged research to develop community capacity, academic researchers worked with certified peer specialists to develop peers research capacity [5]. To prepare certified peer specialists to be involved in research, the principal investigator met with certified peer specialists on the scientific team and also the peer direct service team. Orientation included an open discussion of the current state of the evidence; models for developing behavioral interventions; intervention components; the role of peer interventionists and the peer researcher; and defining expectations of a culture of openness, trust, respect, commitment, flexibility, adaptation, and willingness to compromise. Capacity building included training on the following: (1) research terminology such as pre-posttests, pilot, and outcome measurement; (2) research procedures such as participant safety, informed consent, and data collection; (3) working collaboratively; (4) shared decision-making; and (5) respecting diversity. Instructional methods included experiential learning, role-play, and teach-back method. Academic researchers solicited feedback on how to improve research training.

Once trained, certified peer specialists directed scientific efforts related to the following: (1) text message dose; (2) recruitment—deciding on the location, identifying and hiring, training certified peer specialists using academic profiling, and implementation; (3) identifying outcomes of interest; (4) modified research questions (see below for detailed description); (5) resource allocation—defining equitable pay for certified peer specialists, caseload, and hours required; (6) interpretation of the findings; and (7) dissemination.

The peer direct service team met weekly with the peer supervisor (ie, the peer supervisor was also included in the scientific team) to discuss services delivery, training needs, and modifications to PeerTECH intervention delivery and management procedures. The peer supervisor directed issues to academic researchers weekly during the collaborative research team meetings. Issues brought forth by the peer supervisor included transportation for certified peer specialists, developing a program for matching peers with consumers, and technology training for certified peer specialists. Pragmatic considerations in community-engaged research included flexibility and resource sharing, for example, during training, we included providing financial incentives to certified peer specialists, flexibility with starting times and overestimating timeline, and offering transportation assistance and food.

Ongoing Opportunities for Academic Researchers-Certified Peer Specialists’ Co-Learning

Reciprocal Capacity Building and Learning

We included reciprocal capacity building and co-learning in our application of community-engaged research. On the basis of certified peer specialists’ expertise, peers advocated to examine additional outcomes and also identified potential mechanisms of action. Specifically, peers identified important outcomes including social support, hope, and empowerment. As such, we examined these outcomes. Peers also suggested that without having hope, how is managing ones’ mental and physical health possible? Thus, we defined hope as a mechanism of action on the self-management of medical and psychiatric skill development. As such, academic researchers and certified peer specialists also modified the existing research questions. The original research question included the following: “to what extent does PeerTECH impact self-management skill development?” This research question was modified to examine “to what extent does PeerTECH impact self-management skill development, hope, empowerment, and social support?”

Certified peer specialists worked alongside academic researchers and identified research sites and acceptable screening tools and assisted in hiring, training, and managing peers. The academic researchers-certified peer specialists’ mHealth research group allowed reciprocal capacity building and learning. For example, researchers learned peer history and services delivery practice standards, peer support, and mutuality. The major change this made to the research included modifying the intervention to include peer support in addition to medical and psychiatric self-management skills training, refocusing the intervention from the medical model to the recovery model of services delivery, and the inclusion of consumer-reported measures. In
practice of cultural humility (ie, a process of self-reflection that supports individuals in learning about others’ and their own beliefs and identity [20]), we de-emphasized that researchers were the experts; rather, both groups brought their unique expertise to the team. Cultural humility allowed academic researchers to accept and maintain personal flexibility to allow rapid intervention co-design—consistent with the principles of community-engaged research.

Results

Rapid Iterative Intervention Co-Design

The academic researchers-certified peer specialists’ mHealth research group led the effort to examine the feasibility, acceptability, and preliminary effectiveness of PeerTECH with adults with SMI in a pre-post pilot study. Our study design and findings have been reported elsewhere [21]. PeerTECH showed statistically significant improvements in psychiatric self-management on the illness management and recovery scale (IMRS) [22]. The IMRS is a valid, reliable 15-item scale that assesses domains of illness management [22]. Each item addresses psychiatric illness, management, and recovery. Although we were not powered to detect statistically significant differences, improvements were found including positive changes in self-efficacy for managing health conditions, hope, quality of life, medical self-management skills, and empowerment.

At the pilot study’s conclusion, we conducted a focus group with 3 certified peer specialists involved with PeerTECH and 8 individual interviews with adults with SMI [23]. Both groups agreed technology was a vital component of PeerTECH that allowed health behavior change, self-management therapeutic techniques, engagement in health technology, and peer support [23]. Peers assisted in redesigning the curriculum and requested PeerTECH to promote certified peer specialists’ professional practice standards, including (1) sharing peers’ recovery story, (2) reciprocal learning between peers and consumers whom they support, (3) focusing on the person and their personal goals, and (4) include additional modules on hoarding and trauma-informed care. Certified peer specialists and academic researchers also identified implementation barriers and facilitators to using mHealth in environmental contexts. Barriers identified included text size and the need for the app to be available on a tablet. In response, we are redesigning PeerTECH with the academic researchers-certified peer specialists’ mHealth research group and planning a randomized control trial. In addition, technology allowed everyone to be engaged, regardless of timing and physical barriers.

Engaging Certified Peer Specialists to Cofacilitate Dissemination

Aligned with the community-engaged research principle of activating community assets [5], academic researchers engaged certified peer specialists to cofacilitate dissemination. Certified peer specialists and other stakeholders have been instrumental in disseminating study results through national presentations and peer-reviewed publications. Our research team has historically published and presented with certified peer specialists at national and state conferences. In addition to academic researchers’ dissemination efforts, we utilize other opportunities of dissemination including social media such as Facebook, LinkedIn, and Twitter. To ensure the peers and consumers are informed of our findings, we have worked with certified peer specialists to help us translate the study findings for social media posts to reach a broader audience.

Long-Term Collaboration Between Academic Researchers and Certified Peer Specialists

Aligned with the principle of community-engaged research to commit to long-term collaboration [5], academic researchers and certified peer specialists created a sustainable workgroup. Before the development of the academic researchers-certified peer specialists mHealth research group, the Dartmouth Centers for Health and Aging had no formal research collaborations with certified peer specialists. Although the Dartmouth Centers for Health and Aging has a history of delivering consumer co-led services, certified peer specialists were not involved in the development of these interventions nor were they on research teams. It was essential to work with certified peer specialists and create an infrastructure that would facilitate openness, trust, and respect between academic researchers and certified peer specialists.

Individuals in the academic researchers-certified peer specialist mHealth research group have collaborated on 5 studies over 1 year to advance the role of peers in services delivery using mHealth. We anticipate this group to be long-term as it has secured grant funding from a private foundation to sustain their study. Consistent with principles of community engagement, this approach to community-engaged research has created a rare environment of inclusion through the combination of scientific expertise and certified peer specialists’ expertise (see the Academic Researchers-Certified Peer Specialists mHealth Research Continuum in Multimedia Appendix 1).

Discussion

Principal Findings

This is the first report of a community-engaged research framework that includes certified peer specialists in all stages of research and mHealth intervention development. Unlike the recent models of mHealth community-engaged research that focus on only 1 aspect of community engagement (ie, usability testing), we incorporated peers in all aspects of research and mHealth intervention development. In our framework, peers were equal partners in helping define the problem, creating mHealth intervention content, identifying outcomes of interest, modifying research questions, and identifying research sites, and they also assisted researchers in hiring, training, and managing peers. Finally, peers also guided our dissemination efforts. Our framework can be used as a guide for researchers and certified peer specialists to develop and implement peer-delivered mHealth interventions in community settings.

Conclusions

mHealth intervention development and implementation is a complex scientific process that incorporates multiple disciplines with their own distinctive cultures and expertise [18]. Promoting
empowerment and autonomy with certified peer specialists can be a challenging process in addition to an already complex undertaking. For example, including consumers in mHealth intervention development requires increased time and resources to facilitate equal partnerships, including developing trusting relationships and capacity building; understanding consumers’ opinions, culture, and philosophies; addressing potential relapse; and building and maintaining equal, respectful partnerships. Despite this complex undertaking, we posit that peer-delivered mHealth interventions designed to improve the lives of people with SMI include certified peer specialists with lived experience as experts in every mHealth intervention development and implementation phase.

Acknowledgments
KF’s funding was received from the National Institutes of Mental Health (T32 MH073553-11) and the Brain and Behavior Foundation Early Career Investigator NARSAD award. CG is supported by the National Heart, Lung, and Blood Institute (K23 HL136845). LB is supported by the National Center for Advancing Translational Sciences and the Center for Translational Science Activities (grant number KL2 TR002379), a component of National Institutes of Health (NIH). The views expressed in this study are those of the authors and do not necessarily represent the views of the NIH. The funders had no role in study design, data collection and analysis, the decision to publish, or preparation of the manuscript.

Conflicts of Interest
None declared.

Multimedia Appendix 1
Framework for the Academic Researchers-Certified Peer Specialists mHealth Research Continuum designed to simultaneously inform the development and implementation of a peer-delivered mHealth intervention for adults with serious mental illness.

References


Abbreviations

- I-IMR: integrated illness management and recovery
- IMRS: illness management and recovery scale
- mHealth: mobile health
- NIH: National Institutes of Health
- SMI: serious mental illness
Meaningful Partnerships: Stages of Development of a Patient and Family Advisory Council at a Family Medicine Residency Clinic

Jeffrey D Schlaudecker¹,²*, MD, MEd; Keesha Goodnow¹*, BAE; Anna Goroncy¹,²*, MD; Reid Hartmann¹,²*, MD; Saundra Regan¹*, PhD; Megan Rich¹,²*, MD; Adam Butler¹*; Christopher White¹,²*, MD, JD, MHA

¹Research Division, Department of Family and Community Medicine, University of Cincinnati, Cincinnati, OH, United States
²The Christ Hospital/University of Cincinnati Family Medicine Residency Program, Department of Family and Community Medicine, University of Cincinnati, Cincinnati, OH, United States
*all authors contributed equally

Corresponding Author:
Jeffrey D Schlaudecker, MD, MEd
Research Division
Department of Family and Community Medicine
Medical Sciences Building 4010
231 Albert Sabin Way
Cincinnati, OH, 45267-0504
United States
Phone: 1 513 721 2221
Fax: 1 513 345 6665
Email: jeffrey.schlaudecker@uc.edu

Abstract

Background: Partnering with patients and families is a crucial step in optimizing health. A patient and family advisory council (PFAC) is a group of patients and family members working together collaboratively with providers and staff to improve health care.

Objective: This study aimed to describe the creation of a PFAC within a family medicine residency clinic. To understand the successful development of a PFAC, challenges, potential barriers, and positive outcomes of a meaningful partnership will be reported.

Methods: The stages of PFAC development include leadership team formation and initial training, PFAC member recruitment, and meeting launch. Following a description of each stage, outcomes are outlined and lessons learned are discussed. PFAC members completed an open-ended survey and participated in a focus group interview at the completion of the first year. Interviewees provided feedback regarding (1) favorite aspects or experiences, (2) PFAC impact on a family medicine clinic, and (3) future projects to improve care. Common themes will be presented.

Results: The composition of the PFAC consisted of 18 advisors, including 8 patient and family advisors, 4 staff advisors, 4 resident physician advisors, and 2 faculty physician advisors. The average meeting attendance was 12 members over 11 meetings in the span of the first year. A total of 13 out of 13 (100%) surveyed participants were satisfied with their experience serving on the PFAC.

Conclusions: PFACs provide a platform for patient engagement and an opportunity to drive home key concepts around collaboration within a residency training program. A framework for the creation of a PFAC, along with lessons learned, can be utilized to advise other residency programs in developing and evaluating meaningful PFACs.

(J Participat Med 2019;11(1):e12105) doi:10.2196/12105

KEYWORDS
advisory committee; patient-centered care; patient engagement; graduate medical education

http://jopm.jmir.org/2019/1/e12105/
**Introduction**

**Background**

The core concepts of patient- and family-centered care (PFCC) are based on meaningful partnerships between patients, families, and the health care team. These partnerships promote dignity and respect; encourage information sharing between the provider and patient; foster participation in shared decision making; and cultivate collaboration between the patient, family, and health care team involved in the patient’s care [1]. Collaboration and participation across all ages and locations of care ensure that patients and their families act as included partners in care. Partnering with an active and engaged patient and family generates improved patient satisfaction and delivers upon the goal of higher quality care [2,3]. Providing the authentic voice of patients and families with the health care team is essential in moving health care to a fuller realization of optimized health for all. A patient and family advisory council (PFAC) is a group of patients and family members working together collaboratively with providers and staff to improve health care. A PFAC has shown success in several outcomes including patient experience and satisfaction [4,5]. The Centers for Medicare and Medicaid Services (CMS) recently introduced an innovation in primary care transformation, the Comprehensive Primary Care Initiative Plus (CPC+) [6]. Of the tenets of CPC+, 1 is patient and caregiver engagement, with the establishment of a PFAC being one requirement to fulfill that tenet [7,8].

The Christ Hospital Family Medicine Center is located in Cincinnati, Ohio. A total of 9 faculty providers and 26 resident physicians provide full-spectrum primary care for approximately 7000 patients, with 23,000 annual visits in a practice that is 60% Medicare/Medicaid and the remainder private insurance. The office was certified as a patient-centered medical home in 2010.

**Objectives**

This study describes 1 program’s experience launching a PFAC within a family medicine residency clinic. We discuss the stages of PFAC development, which include PFAC leadership team formation and initial training, member recruitment, and meeting launch. PFAC evaluation included an open-ended survey and focus group interview at the completion of the first year.

**Methods**

**Patient and Family Advisory Council Leadership Team Formation and Initial Training**

A desire for family medicine resident physicians to engage with patients in office-based practice improvement initiatives was broadcast to faculty stakeholders, including the program director and clinic medical director. These faculty stakeholders agreed that a PFAC could be an ideal educational tool at our residency program. Requirements of the CMS’s 2016 CPC+ program of primary care payment restructuring to include a PFAC additionally piqued the interest of the health system and clinic medical director. A project manager was hired to manage the day-to-day operations of a related grant, and this person additionally served as the PFAC coordinator. Following approval of the residency program director and clinic medical director, the PFAC leadership team was formed. This team included several authors of this paper, including 2 residency faculty members and the PFAC coordinator.

Initial steps of the PFAC leadership team focused on identifying key practice stakeholders including the clinic nurse manager. Exploratory meetings were held with clinic and residency leadership to bolster agreement on the goal of the PFAC. The PFAC would be utilized to introduce patient voice into clinic issues, including residency-based clinic quality improvement projects. In addition, our PFAC would serve as a training model for residents to learn about the benefits of meaningful collaboration in their future practices.

To attain knowledge about the development and implementation of a PFAC, the PFAC leadership team attended a national 3-day seminar hosted by The Institute for Patient and Family Centered Care [1]. The seminar was a platform to learn from experts fostering genuine partnerships with patients, family members, and health care teams to improve the safety and quality in health care. Following this initial training, the team completed a formal action plan addressing the next steps needed to recruit and launch a PFAC.

**Patient and Family Advisory Council Member Recruitment**

On the basis of best-practice examples, it is critical to identify a representative sample of patients and family members who have a wide variety of clinic experiences and reflect the diversity of the patient population. It is not necessary for all advisors to have universally positive experiences with the health care system or clinic; however, all PFAC members should have a desire to see the clinic improve. Outreach methods to enlist advisors included brochures and posters. A pocket information card we developed was our most effective recruitment strategy. Our providers carried this business card-sized tool in the clinic, which allowed them to invite engaged patients or family members in real time to consider joining our PFAC during the clinic visit.

Once patient or family members were identified as potential PFAC members, the medical director would call prospective candidates to personally invite them and validate the partnership between provider and patient. This point of contact also served to obtain permission for the PFAC coordinator, who was not an employee of the organization owning the clinic, to call to further discuss the PFAC. The PFAC coordinator then called each potential PFAC member to provide detailed information and conducted a short phone interview. The aim of this phone call was to communicate the purpose of the PFAC, describe the role of advisors, set expectations for involvement, and query potential barriers to attending meetings. In addition, patients and family members shared personal information and experiences that were helpful in guiding PFAC design. During the phone interview, the PFAC coordinator also obtained motivation and rationale for involvement in the PFAC. Although we have not barred a patient or family member from membership, we recognize that having an awareness of personal agendas is important to ensure that there is no conflict between patient or family member motivation and PFAC goals. At the completion of the call, an application was sent to potential PFAC members.
members to collect contact information and preferences regarding meeting time.

Our PFAC membership included staff and physician members as well as patients and family members. Resident participation as PFAC members was solicited via email; 4 residents expressed interest and enthusiasm for participating. To maximize attendance, the PFAC leadership team decided on a monthly meeting schedule held from 5:00 pm to 6:00 pm on Wednesday evenings. The PFAC leadership team developed a curriculum for introducing health care–related topics, such as Health Insurance Portability and Accountability Act and quality improvement strategies, to the new PFAC members. A timeline for the process is depicted in Figure 1.

In accordance with hospital requirements, all volunteers, including PFAC advisors, completed a background check. As advisors occasionally encounter other patients and family members while attending meetings, it was important to follow this protocol to protect patient safety and privacy. Each PFAC member was asked to commit to a 1-year term with the option to re-enroll at the end of the term. This was critical in building a sense of community and trust for the partnership to be effective, while also providing a time-limited commitment that acknowledges busy lives. To support the advisors in keeping this commitment, free childcare was provided on site. Neither office staff nor PFAC advisors were reimbursed for their time, although snacks were provided at every meeting. The final composition of the PFAC was 18 PFAC members, including 8 patient/family advisors, 4 staff advisors, 4 resident advisors, and 2 faculty advisors.

Figure 1. Patient and family advisory council creation timeline. PFAC: patient and family advisory council; QI: quality improvement.

Figure 2. Patient and family advisory council sample meeting agenda. PDSA: Plan-Do-Study-Act.
Patient and Family Advisory Council Meeting Launch

The initial meeting of the PFAC took place in fall 2016 and focused on key concepts for PFAC members, including basic health care concepts and confidentiality and communication strategies. Threaded throughout subsequent meetings were team-building exercises to build a sense of community, opportunities to practice sharing important health care–related stories, and discussion of current and relevant clinic issues to elicit immediate feedback from advisors. The basic tenants of clinic-based quality improvement, including tools like Plan-Do-Study-Act (PDSA) cycles, were also introduced, as were occasions to examine the office space and to identify areas for improvement projects (Figure 2).

PFAC recommendations helped identify the final projects: waiting room and exam room redesign to improve the patient experience. The final meetings focused on completing PDSA cycles including pre- and postintervention data collection and analysis (Figure 3).

Measurement of Patient and Family Advisory Council Impact

To evaluate the PFAC implementation, we used a mixed-methods approach. We tracked the number of meetings held, number of attendees at each meeting, and roles of those who attended and conducted a focus group interview of the PFAC members at the end of the year to evaluate impressions, lessons learned, and suggestions for subsequent years. All PFAC members also completed an open-ended survey so that they could evaluate the year anonymously [9]. The evaluation team, consisting of a qualitative social science researcher, family physician, project manager, and premed student, analyzed data from open-ended surveys and focus group interviews, generating descriptive codes to create consensus themes.

Figure 3. Patient and family advisory council monthly meeting highlights. HIPAA: Health Insurance Portability and Accountability Act; PDSA: Plan-Do-Study-Act; PFCC: patient- and family-centered care; QI: quality improvement.
Results

Group Composition
The composition of the PFAC consisted of 18 advisors, including 8 patient/family advisors, 4 staff advisors, 4 resident advisors, and 2 faculty advisors. The average meeting attendance was 12 members over 11 meetings in the span of the first year, with the most attended meeting having 15 members and the least attended having 9 PFAC members in attendance. Key leadership was present at every meeting, including the medical director and office nurse manager. Despite our best efforts, there was some attrition of patient and family advisors. By year’s end, there were 4 patient/family advisors regularly attending meetings, down from our 8 initial recruits.

Qualitative Results
The PFAC’s first year concluded with feedback from the participants focusing on what they considered favorite aspects or experiences, impressions on their impact on the family medicine center, and guidance on future projects to improve patient care (Table 1). There were 13 respondents, including 7 patient and family advisors, 2 staff advisors, and 4 resident/physician advisors.

Table 1. Qualitative data from the patient and family advisory council group.

<table>
<thead>
<tr>
<th>Topic and theme</th>
<th>Quote</th>
</tr>
</thead>
<tbody>
<tr>
<td>Favorite aspects or experiences</td>
<td></td>
</tr>
<tr>
<td>Walkabout</td>
<td>That was the first time I had ever spent a significant amount of time in our waiting room, where it was before and just sitting down in one of those chairs it was like, Whaa, this gives me some perspective. [Physician PFAC member]</td>
</tr>
<tr>
<td></td>
<td>I really enjoyed the walkabout and the conversations that followed. It felt like a really productive use of time and energy. [Patient PFAC member]</td>
</tr>
<tr>
<td>Sharing your story</td>
<td>Sharing personal moments/stories at start of meeting helped understand where each member’s perspective is built from. Also bond as a group. [Patient PFAC member]</td>
</tr>
<tr>
<td></td>
<td>...sharing stories moment, in a safe place, talk about what it was like for my sister to just have walked through this health crisis and the thing that hurt the most was the thing the Medical Assistant said off-hand rooming her into the office. Just talk through these sorts of things. [Resident PFAC member]</td>
</tr>
<tr>
<td>PFAC involvement in practice meeting</td>
<td>It was powerful to take PFAC concerns back to the FMC meeting and have that steer our efforts in QI development. The collective knowledge and goals from the group provide focus and force to getting projects accomplished, much more than any individual effort or goals could. [Physician PFAC member]</td>
</tr>
<tr>
<td>PFAC’s most positive impact on the family medicine center</td>
<td>The PFAC’s very existence and its positive reception by the medical professional caregivers are the sources of any impact on care delivered at the Fam. Practice Center. [Patient PFAC member]</td>
</tr>
<tr>
<td></td>
<td>Developing open communication lines between providers and patients. [Patient PFAC member]</td>
</tr>
<tr>
<td></td>
<td>Incorporating patient voices into the development of new projects. Working to make it a more patient-centered culture. [Patient PFAC member]</td>
</tr>
<tr>
<td>Future projects to improve patient care: communication</td>
<td></td>
</tr>
<tr>
<td>Phone</td>
<td>I feel like something that is frustrating for me at times is when I call and have a question, whether it’s for a doctor or a nurse, and I kind of feel like it goes into a black hole, and I don’t get a response or it takes weeks. I would just love to see that process streamlined or figured out. [Patient PFAC member]</td>
</tr>
<tr>
<td>MyChart</td>
<td>I think there is a breakdown of communication between patients, doctors, phone staff, and MyChart. [Patient PFAC member]</td>
</tr>
<tr>
<td>Waiting time</td>
<td>Perhaps have updates for patients who have been roomed for &gt;15 minutes. An update on where the provider is on the schedule may be helpful! [Patient PFAC member]</td>
</tr>
</tbody>
</table>

The topics identified common areas of the PFAC that members found to be positive and influential, indicating that it may be appropriate to continue to implement the corresponding experiences into the council’s regular operation. The walkabout was an important activity to experience the clinic through the patient lens and identify needs of the practice. Sharing stories is integral in identifying blind spots and providing context to know what needs to be improved. The PFAC involvement in practice meetings helped emphasize the patient and family voice and validate this partnership. The PFAC’s positive impact on the family medicine center was a result of building a vehicle for collaboration to improve the patient experience. Overall, 13 out of 13 (100%) advisors that completed the surveys were satisfied with their experience serving on the PFAC, with 1 member commenting:

It’s been great to be a part of something that is helping make a tangible change.

A common theme throughout centered on the value of member participation. This placed a focal point on ways in which PFAC members, particularly the patient and family advisors, can take a leading role in generating new ideas for the council to address in further driving the PFAC forward for the coming years.

aPFAC: patient and family advisory council.
Developing a safe environment with a strong sense of community and trust was the foundation for advisor engagement, promoting crucial input for relevant quality improvement ideas and ownership of projects. In analyzing future projects suggested by PFAC advisors, communication is at the core of each theme, a cornerstone of PFCC.

Discussion

Lessons Learned to Guide the Patient and Family Advisory Council’s Future

Our approach to PFAC implementation started with clinic and residency program leadership and garnered support before involving office staff or resident learners. Although leadership approval is certainly an important step early in the process, a top-down approach led to some limitations. We found that very early in the process, resident physicians and office staff seemed less invested in the PFAC and did not always recognize how collaboration with patient and family advisors could enhance the quality of care. Once up and running, investment was high, and in future iterations, we plan to take an interprofessional approach to involve leadership, physicians, residents, staff, and patients/family members from initial conception. Having representation from all stakeholders during the creation phase may prove beneficial to determine shared goals and promote support. The literature supports councils seeking to attain buy-in early to promote success [10].

The attrition rate of patient and family advisors was disappointing but not unexpected. There are many barriers to our patients attending meetings, from transportation problems to jobs with evening shifts to childcare. We were able to address some issues (such as childcare) but not all. The literature suggests most PFACs find it problematic to establish a diverse membership [11]. Although demographics in our office reflect the city composition with 50% patients of color, our PFAC had only 2 non-white members. For the upcoming cycle, we are recruiting more patient and family advisors, with focused representation from all stakeholders during the creation phase may prove beneficial to determine shared goals and promote support. The literature supports councils seeking to attain buy-in early to promote success [10].

The literature highlights the need for an administrative manager to help make the PFAC more valuable and productive [11]. To keep attendance high, the PFAC coordinator kept in close contact with patient and family advisors through email, text, and phone calls based on advisors’ preferred forms of communication. This included reminders about upcoming meetings, sharing of “homework” assignments, and troubleshooting any potential conflicts that could interfere with meeting attendance. The estimated time required for coordinator oversight, not including work related to the collection of metrics related to the PFAC evaluation, was on average 10 hours per month. Time fluctuated throughout the year depending on the needs of the PFAC. Recruitment, implementation of projects, and year-end evaluation required additional time and effort.

Finally, as a residency training site with 26 learners, physician faculty are exploring ways to seamlessly incorporate PFAC concepts into the general residency teaching curriculum. Our PFAC is not only an ideal driver in identifying quality improvement projects but also serves as a critical partner in resident training. We have focused on harnessing patient and family advisors as teachers by inviting them to share at resident conferences. In addition, we now have patient and family advisors attend our practice meetings to promote buy-in, strengthen communication, and enhance their role as teachers. Although 3 to 4 resident physician PFAC members were felt to be ideally representative, the PFAC leadership team has also chosen to rotate residents every year to give as many residents as possible the chance to experience the PFAC. Though it is not possible for all residents to sit on the PFAC, more exposure to the patient and family advisors and their stories and perspectives can assist in driving home key concepts around patient participation and collaboration.

Conclusions

The health care industry now recognizes that partnering with patients and families is a crucial step in optimizing health. This trend marks an evolutionary shift from the paternalistic practices of the past toward a new model of health care partnership between providers, patients, and families. Our residency-based family medicine center launched a PFAC in the fall of 2016 to collaborate with patients and family members beyond the traditional patient encounter. Our initial successes and lessons learned have cemented the belief in the positive outcomes possible with meaningful partnerships with patients and families.

Acknowledgments

This work was supported by the Health Resources and Services Administration Primary Care Training and Enhancement Grant T0BHP28567 (Partnering with Underserved Patients: A Novel Health Transformation Curriculum).

Conflicts of Interest

None declared.

Multimedia Appendix 1

Video abstract.

[MP4 File (MP4 Video), 69MB - jopm_v11i1e12105_app1.mp4 ]

References


Abbreviations

CMS: Centers for Medicare and Medicaid Services
CPC+: Comprehensive Primary Care Initiative Plus
PDSA: Plan-Do-Study-Act
PFAC: patient and family advisory council
PFCC: patient- and family-centered care

©Jeffrey D Schlaudecker, Keesha Goodnow, Anna Goroncy, Reid Hartmann, Saundra Regan, Megan Rich, Adam Butler, Christopher White. Originally published in Journal of Participatory Medicine (http://jopm.jmir.org), 20.03.2019. This is an open-access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work, first published in Journal of Participatory Medicine, is properly cited. The complete bibliographic information, a link to the original publication on http://jopm.jmir.org, as well as this copyright and license information must be included.