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Monitoring and Evaluation of Patient Engagement in Health Product Research and Development: Co-Creating a Framework for Community Advisory Boards

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Purpose While patient engagement is becoming more customary in developing health products, its monitoring and evaluation to understand processes and enhance impact are challenging. This article describes a patient engagement monitoring and evaluation (PEME) framework, co-created and tailored to the context of community advisory boards (CABs) for rare diseases in Europe. It can be used to stimulate learning and evaluate impacts of engagement activities.

Methods A participatory approach was used in which data collection and analysis were iterative. The process was based on the principles of interactive learning and action and guided by the PEME framework. Data were collected via document analysis, reflection sessions, a questionnaire, and a workshop.

Results The tailored framework consists of a theory of change model with metrics explaining how CABs can reach their objectives. Of 61 identified metrics, 17 metrics for monitoring the patient engagement process and short-term outcomes were selected, and a “menu” for evaluating long-term impacts was created. Example metrics include “Industry representatives’ understanding of patients’ unmet needs;” “Feeling of trust between stakeholders;” and “Feeling of preparedness.” “Alignment of research programs with patients’ needs” was the highest-ranked metric for long-term impact.

Conclusions Findings suggest that process and short-term outcome metrics could be standardized across CABs, whereas long-term impact metrics may need to be tailored to the collaboration from a proposed menu. Accordingly, we recommend that others adapt and refine the PEME framework as appropriate. The next steps include implementing and testing the evaluation framework to stimulate learning and share impacts. (J Patient Cent Res Rev. 2022;9:46-57.)

Keywords patient engagement; patient participation; rare diseases; program evaluation; drug development

Patient engagement is increasingly integrated along the life cycle of research and development (R&D) for health products (ie, medicines, technologies and services) as an essential element in moving toward a more patient-centered R&D system.1–3 Patient engagement is defined as “the active, meaningful, and collaborative interaction between patients and researchers across all stages” of health product R&D, “where decision-making is guided by patients’ contributions as partners, recognising their specific experiences, values, and expertise.”4 Such interactions have rapidly increased in the last decade, taking many shapes and forms.1,5 However, little is known about how these interactions affect health product R&D processes and their stakeholders.6,7 This knowledge gap makes it difficult to understand best practices and the value of patient engagement, increasing the risk of tokenism and hindering successful embedding of practices in R&D processes.6,8 The need for monitoring and evaluation (M&E) to elucidate whether, how, and why the objectives of patient engagement are achieved has been repeatedly articulated.7,10
Despite the need, M&E of patient engagement initiatives in health product R&D is scarce, and current evidence is often considered anecdotal. Previous literature reviews discussing M&E tools and frameworks report concerns regarding their scientific rigor, lack of co-creation and impact assessment, and limited transferability to other settings. Various challenges underlie these limitations. First, patient engagement initiatives are highly contextualized and relational in nature, and it often takes years before impact can be seen. Consequently, many different factors influence the causal chains leading to impact, impeding the attribution of causality and the generalization of conclusions. Moreover, stakeholders have different needs and priorities, and therefore views vary on what constitutes successful patient engagement.

Partly because of this, there is no consensus on relevant evaluation criteria and tools. While some seek standardization across the field to enable generalization and comparison, others argue that M&E frameworks and tools provide more meaningful insights when tailored to the initiative and co-created with those involved.

This article presents a tailored M&E framework for community advisory boards (CABs) participating in the European Organisation for Rare Diseases (EURODIS) program (collectively known as EuroCAB). CABs are autonomous groups of patient representatives (ie, CAB members), nominated and selected by their community, who offer their expertise to public and private sponsors of research in their disease area (mainly, the biopharmaceutical industry) through advisory meetings. They aim to align clinical research to community needs and realities. EuroCAB was established to provide training and support for CABs in the field of rare diseases. CAB members and industry representatives expressed the need for evaluation to gain a deeper understanding of the CAB processes and outcomes in order to identify strategies to enhance and demonstrate their impact. Accordingly, the objective of this study is to develop a meaningful and feasible M&E framework for these CABs.

**METHODS**

**Theoretical Background**

The development of a framework for CABs is supported by the patient engagement monitoring and evaluation (PEME) framework, which can be tailored — in co-creation with all stakeholders — to a specific initiative. The starting point is developing a theory of change model to collaboratively explain the steps toward the overarching objectives of the initiative, the relationships between these steps, and the underlying assumptions and context factors of how and why an initiative works. Iteratively, quantitative or qualitative metrics (sometimes referred to as “indicators” or “measures”) linked to the steps of the theory of change are identified and prioritized. Through discussion, the processes influencing success are described and illustrated, and consensus may be reached on a visual narrative model of the steps toward impact, accompanied by relevant metrics.

The model and metrics form the tailored framework, which is structured along 6 components: 1) objectives, 2) input, 3) activities/process, 4) learnings and changes, 5) impacts, and 6) context (Figure 1). Hence, the PEME framework focuses on generative causation to ensure that M&E stimulates reflection on how and why impacts are achieved, thereby inspiring change.

**Setting**

The study was conducted as part of the Patients Active in Research And Dialogue for an Improved Generation of Medicines (PARADIGM) consortium. The EuroCAB program was one of the case studies in this consortium. Currently, 5 CABs participate in the program: Duchenne Muscular Dystrophy, Cystic Fibrosis, Cystinosis, Lymphomas, and Hereditary Haemorrhagic Telangiectasia. CAB members and industry representatives collaboratively organize the CAB meetings, which strive for sustainable partnership that influences R&D decision-making by focusing on collective thinking and knowledge exchange between patient communities and researchers. The EuroCAB program applied and tailored a preliminary version of the PEME framework to develop an M&E framework for CABs in the rare disease context.

**Participatory Approach**

A participatory approach was adopted for framework development. This process was based on the principles of interactive learning and action and comprised 4 iterative data collection and analysis phases: 1) initiation and preparation; 2) collection, exchange, and integration of information; 3) priority setting; and 4) validation and planning (Table 1). Research activities took place between January 2019 and May 2020. The project team (authors S.E.F. and L.E.V., who gathered and analyzed the data, and R.C. and F.H., who guided the process and interpreted findings) held regular meetings to reflect on the findings, refine the framework, and plan next steps. Online Appendix A provides an overview of terms and definitions used throughout this study.

Vrije Universiteit Amsterdam’s ethics review committee determined that the research proposal complied with the ethical guidelines of its Faculty of Science.

**Recruitment, and Data Collection/Analysis**

**Phase 1 – Initiation and Preparation:** In phase 1, the project team aimed to understand the context of the
EuroCAB program and reviewed the existing M&E practices to develop a preliminary case description. This was based on document review (CAB description and agreements, meeting minutes, and feedback tools developed by EUORDIS) and informal discussions with the project team.

**Phase 2 – Collection, Exchange, and Integration of Information:** In phase 2, reflection sessions with CAB members and industry representatives were held to reflect on their objectives, collaboration processes, and experiences with M&E. The aim of this phase was to identify meaningful metrics to iteratively develop and refine a preliminary M&E framework. Participants were invited to take part via the EuroCAB coordinator (R.C.) using a convenience sampling method. Participation was voluntary. Two group reflection sessions (~2 hours) were held, one with Duchenne Muscular Dystrophy CAB members (n=11) and one with Cystic Fibrosis CAB members (n=11). In addition, one-to-one reflection conversations (~1 hour) were conducted with 4 industry representatives from different companies and 2 CAB chairs (from Duchenne Muscular Dystrophy CAB and Lymphomas CAB). The preliminary framework and preliminary findings from the feedback tools developed by EUORDIS were presented and discussed during a workshop at a EUORDIS event with patients and companies. Notes were taken for all sessions.

**Phases 1 and 2 – Analysis:** During data collection, the theory of change was drafted and iteratively refined to develop a detailed representation of the CAB collaborations. Using the software program ATLAS.ti (Scientific Software Development GmbH), documents and notes from phases 1 and 2 were first analyzed deductively by author S.E.F. based on components of the PEME framework. Objectives and metrics within these components were identified via an inductive approach of open coding (identifying and describing insights), followed by axial coding (creating subcategories) by S.E.F. and L.E.V. Identified metrics were linked to the steps in the theory of change, resulting in a preliminary framework.

**Phase 3 – Prioritization:** Phase 3 aimed to prioritize the objectives and metrics identified in phase 2 to develop a set of meaningful metrics through a priority-setting questionnaire consisting of three parts: 1) role and experiences; 2) selection of applicable objectives and ranking of those selected; and 3) selection of up to 5 most relevant metrics across the core components of PEME (input, activities/process, learnings and changes, and impact). The impact metrics were divided into short- and long-term impact based on when they could be measured — before or after the start of the discussed study. The questionnaire was reviewed by R.C., F.H., and one CAB chair. Industry representatives and members of all CABs were invited to participate in the questionnaire via the CAB chairs or the CAB coordinator. Participation
was voluntary. Of the 30 respondents who completed part 1 and 2 of the questionnaire (Table 2), 27 also completed part 3. Individual responses were anonymous and kept confidential.

**Phase 3 – Analysis:** Data from the questionnaire were analyzed via descriptive statistics using Microsoft Excel. The weighted average ranking of the objectives was calculated (rank 1 was assigned a weight of 7, etc). Unranked objectives were assigned a weight of 0. Metrics were ranked based on how often they were selected, resulting in a prioritized list of all metrics. Respondents could not indicate the relative weight for each of their priority metrics, so to do justice to the gross prioritization, clusters of high-, medium-, and low-priority metrics were created. Trends in prioritization of metrics among stakeholder roles, disease areas, and number of meetings attended were examined. Since industry representatives had only engaged with the Cystic Fibrosis CAB, their responses were excluded from the stratification by disease area to avoid potential bias.

**Phase 4 – Validation and Planning:** In phase 4, a digital dialogue workshop was held to validate the theory of change and set of metrics for M&E and gather ideas for data collection and reporting. The 5 CAB chairs and 5 industry representatives were invited. Three CAB chairs participated, 2 could not. A summary of the questionnaire results and the drafted tailored framework were shared with participants in advance of the workshop. Using polling tools, participants were asked to what extent they agreed with the tailored framework and the “high-priority” metrics and about what reporting structures they preferred, followed by discussion. Postworkshop,
detailed notes were used by the project team and co-authors of this article to make final adjustments to the tailored framework.

RESULTS

The participatory approach resulted in a tailored framework for the EuroCAB program, including a theory of change explaining how and why objectives are reached, 17 high-priority metrics (out of 61 identified), and a menu of possible long-term impact metrics. Herein, we describe each part of the tailored framework, including rationales for metric selection.

Objectives

Seven objectives for the CABs were identified. The prioritization questionnaire showed that, on average, CAB members and industry representatives considered the objective “to ensure products better meet patients’ unmet medical needs” as the most important (Figure 2). The relative importance of the other objectives differed somewhat by stakeholder group. For example, industry ranked “enhance transparency and trust” higher than did CAB members, whereas CAB members found “diminish access inequalities” and “improve efficiency and speed” more important. There were also small differences between disease areas. For example, the Cystic Fibrosis CAB found “diminish access inequalities” more important than other CABs did.

Theory of Change

Starting from the objectives, the actions needed to reach them, underlying assumptions, and context factors were iteratively mapped to develop a theory of change (Online Appendix B). Figure 3 shows an example of a theoretical pathway. The underlying assumptions describe the conditions that stakeholders deemed necessary for the proposed steps to take place. Also, several contextual factors influencing whether objectives of an initiative are achieved were identified and indicated. Each step in the theory of change can be evaluated with metrics (indicated with codes referring to the metrics in Table 3). In Table 3, data collection methods per metric are suggested. The following section highlights some rationales for selecting metrics and describe the links between them.

Metrics

The metric “Feeling of trust between stakeholders” was selected most often. This metric was categorized under activities/process and impact because while some indicated that improved trust is itself an impact, others felt that it is more a means to an end, strongly related to the metric “Feeling of openness and transparency.” In phase 2, most participants indicated that trust is a precondition for the sustainable long-term collaborations and that it takes time to develop:

“The more often you meet a company, you can tell that they have the feeling that we have important things to say, and that is exactly what we are trying to give them. They listen, they respect … they are very, very respectful of our opinion and how we interact with them, and that has led to the trust that is absolutely vital if you are going into confidential information … and it takes time for a relationship like that to really materialize.” (CAB chair 1)

The learnings metrics “Industry’s understanding of patients’ unmet needs,” “Degree to which the meeting helped to inform research goals,” and “Industry’s understanding of patient-relevant outcomes measures” were considered indicative of whether patient needs and experiences are taken into account in R&D processes, aligning closely to the most important objective. The metric “CAB members’ understanding of decision-making in health product R&D” was selected less often. Nevertheless, this metric was included because mutual knowledge exchange is considered a key principle of the CABs and should therefore be assessed from both perspectives.

Table 2. Respondents to Questionnaire in Priority-Setting Phase

<table>
<thead>
<tr>
<th>Demographics</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stakeholder group</td>
<td></td>
</tr>
<tr>
<td>CAB member</td>
<td>24</td>
</tr>
<tr>
<td>Industry representative</td>
<td>6*</td>
</tr>
<tr>
<td>Total</td>
<td>30</td>
</tr>
<tr>
<td>Disease-specific CAB</td>
<td>7</td>
</tr>
<tr>
<td>Cystic Fibrosis</td>
<td>7</td>
</tr>
<tr>
<td>Duchenne Muscular Dystrophy</td>
<td>6</td>
</tr>
<tr>
<td>Cystinosis</td>
<td>4</td>
</tr>
<tr>
<td>Hereditary Haemorrhagic Telangiectasia</td>
<td>7</td>
</tr>
<tr>
<td>Total</td>
<td>24</td>
</tr>
<tr>
<td>Number of CAB meetings attended</td>
<td></td>
</tr>
<tr>
<td>None (in the planning phase)</td>
<td>6</td>
</tr>
<tr>
<td>≥1</td>
<td>24</td>
</tr>
<tr>
<td>Total</td>
<td>30</td>
</tr>
</tbody>
</table>

*27 of 30 respondents completed part 3 of the questionnaire (selection of metrics).

Industry representative respondents all collaborated with the Cystic Fibrosis CAB.

Since all industry representative respondents worked with the Cystic Fibrosis CAB, their responses were excluded from stratification per disease area.

CAB, community advisory board.
To improve industry’s understanding of patient needs and reach the objectives, it was often emphasized that the “right” people need to attend the meeting. Therefore, the metric “Diversity of stakeholders” was included. Diversity of CAB members ensures that the insights and recommendations reflect to some extent those of the overall patient population. Diversity of industry representatives (e.g., in seniority level) indicates the scope of the influence of CAB members on decision-making, as participants noted that broad diversity helps to spread awareness and understanding of patient needs throughout the company.

The metric “Degree to which the collaboration helps to demonstrate the value of a product to regulators and health technology assessment bodies” was selected to capture whether industry representatives report an influence of the collaboration on decision-making beyond their company. Moreover, the metrics “Degree to which outcome measures of a trial are patient-centric” and “Expected study participant burden” were selected to assess whether a trial is well designed, both in terms of the quality of evidence generated and participant experience, as these topics are often discussed in the meetings and patient-centric design is important to companies seeking to recruit quickly, minimize and prevent study drop-outs, and finish trials on time. One participant emphasized that, compared to longer-term impact metrics, the expected burden on study participants could be relatively easily attributed to patient engagement, for example, by reducing the number of hospital visits in a trial. Therefore, this metric was thought to provide compelling evidence of the value of engaging patients.

To track whether the changes needed to reach the impacts are made, more operational metrics, such as “Number and type of planned or recommended actions and changes (not) implemented” were selected. Tracking follow-up after the meeting is essential to ensure progress and prevent repetition:

“In the end, we completely lost track of everything because people were not in charge of the same things. And if there are no clear action steps, and there weren’t any because of the lack of alignment on the topics discussed, then it is very challenging [to make progress].” (CAB chair 3)

Several CAB members pointed out that they would like to receive more feedback from industry on where change is (not) happening to help them understand potential limitations industry may have and reprioritize discussion...
Figure 3. Example of a pathway of the theory of change for community advisory boards in the EuroCAB program. Bright color boxes: actions needed to achieve the objectives. Faded color boxes: underlying assumptions (conditions deemed necessary for the steps to take place). Light gray box: context factors (external influences). (A2), (I3), (L5), etc link to that respective metric defined in Table 3. For the full theory of change, see Online Appendix B. CAB, community advisory board; CM, CAB member; HTA, health technology assessment; IR, industry representative; R&D, research and development.
topics. Likewise, industry representatives indicated that an open discussion on expectations helps to align concrete goals and timeframes and prevent disappointment and frustration:

“It [tracking priorities, discussions, and actions] can also support the discussion around is it realistic that an impact can happen on this priority. ... If you see that you keep on hitting a block on certain type of topic you can just say: ‘Okay, we agree that we are not going to make progress here so we can just kind of focus on other topics.’” (Industry representative 1)

Accordingly, the metric “Stakeholders’ priorities for the meeting” was selected because it could indicate whether the most important topics are discussed. True co-creation of the agenda and thorough preparation were considered...
essential because these create belief in the value and ownership of engagement among all involved.

It was suggested to exclude the long-term impact metrics from the set and include them instead in the tailored framework via a “menu” approach, as presented in Table 4. Participants indicated that long-term impact metrics are valuable as aspirational targets, but many contextual influences may make it difficult to attribute causality. Measuring long-term impact would therefore require thorough and continuous tracking of decisions and changes. The menu allows stakeholders to prioritize, through dialogue, the most meaningful metric(s) for their collaboration while ensuring the burden of measurement remains feasible.

**DISCUSSION**

In this study, the preliminary PEME framework was tailored in co-creation with relevant stakeholders to the context of CABs in the EuroCAB program. The various information needs shaped the tailored framework. Each collaboration between patients and industry is unique, and tailoring M&E by respective context can improve the relevance of the information collected, enhancing learning and reflexivity. Such continuous reflection is needed to understand under what conditions progress toward the objectives is made and to develop strategies to enhance impact. On the other hand, standardized M&E enables data aggregation, generalization, and comparison with other engagement initiatives, thereby better meeting accountability purposes. We aimed to balance these purposes by tailoring a framework to the CAB method and rare diseases context while also standardizing a set of metrics across all CABs in the EuroCAB program, combining metrics for learning and accountability. Our results suggest that “input,” “activities/process,” “learnings and changes,” and “short-term impact” metrics could be standardized across CABs, while the long-term impact metrics could best be tailored to the objectives, capacity, and resources of the collaboration. This resulted in a combination of fixed and flexible sets of metrics.

In line with earlier studies, the results suggest that stakeholders’ M&E priorities differ because of varying interests and needs. In alignment with the CABs’ focus on equal partnership and mutual knowledge exchange, the framework and metrics were constructed from the perspectives both of CAB members and industry and will be assessed from both perspectives when possible. The recognition that the metrics need to be meaningful for all stakeholders underlines the importance of tailoring M&E to enhance impact. Some of the selected metrics are in line with other studies, such as preparedness, diversity of patients, the recommendations (not) implemented and why, and the changes in research questions and design. These metrics might be applicable to patient engagement initiatives in various contexts. Other metrics may be particularly relevant in the rare disease context, such as “Number of patients in preapproval access programs.”

**Table 4. “Menu” of Long-Term Impact Metrics for EuroCABs**

<table>
<thead>
<tr>
<th>Long-term impact metrics</th>
<th>Number of stakeholders who selected metric (N=27)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Perceptions on the alignment of research programs with patients’ needs</td>
<td>16</td>
</tr>
<tr>
<td>Time until regulatory approval and health technology assessment reimbursement decisions</td>
<td>14</td>
</tr>
<tr>
<td>Retention rate (number/percentage of participants retained in a trial and assessed)</td>
<td>14</td>
</tr>
<tr>
<td>Patient satisfaction with participation in a trial</td>
<td>13</td>
</tr>
<tr>
<td>Quality of a product (eg, drug, technology, service, information materials)</td>
<td>12</td>
</tr>
<tr>
<td>Time from start until the end of a trial (first participant in/last participant out)</td>
<td>12</td>
</tr>
<tr>
<td>Number of avoidable amendments (often costly changes to a trial protocol as a result of operational barriers)</td>
<td>11</td>
</tr>
<tr>
<td>Number of patients in a preapproval access program</td>
<td>11</td>
</tr>
<tr>
<td>Diversity of participants in a trial (demographics, disease subtype, hard to reach populations, etc)</td>
<td>11</td>
</tr>
<tr>
<td>Number of patients who have access to clinical trials</td>
<td>8</td>
</tr>
<tr>
<td>Recruitment rate (number of participants recruited per site per month)</td>
<td>7</td>
</tr>
<tr>
<td>Reputation of industry within the patient community</td>
<td>6</td>
</tr>
</tbody>
</table>

*Long-term impact metrics were not coded and indicated in the theory of change, as no consensus was reached and no methods were discussed.*
because of the high unmet medical need. Future studies could further explore the framework’s generalizability to other disease contexts outside of rare diseases.

The PEME framework supported the identification of relevant metrics and ensured that the tailored framework comprehensively captures the relationship between input and impacts. This improves understanding of the processes that influence success and can enable the attribution of causality. However, it remains difficult to prioritize metrics. The selected metrics do not fully reflect the potential value of the CABs, as they focus mainly on R&D while discussed topics and impacts may be broader. Despite excluding many steps of the theory of change from assessment, the set of metrics is still ambitious and implementation may prove challenging. This difficulty in prioritization could perhaps be overcome by urging participants to make their selection criteria more explicit (eg, relevant, feasible, transferable). It is important to note that the tailored framework is meant as a flexible tool for reflection — refined by each CAB/industry relationship — to enhance the collaboration processes. Next steps include developing and refining quantitative and qualitative data collection methods and implementing the evaluation framework to encourage learning and share impacts.

Strengths and Limitations
During the process, we aimed to integrate stakeholders’ different perspectives while respecting diverse viewpoints through democratic participation complemented by discussions on the results. The collaborative approach created a common language and increased ownership of the tailored framework, which we see as a strength. It also supported the “relational work” of aligning expectations and developing a shared purpose. Thus, this process is itself an extension of the collaborative nature of the EuroCAB program.

A limitation was that CAB members and industry representatives of each of the 5 CABs could not participate in all phases, and the balance between CAB members and industry was not always exactly equal. However, all phases included a mix of perspectives. Also, the framework’s linear success-oriented structure may be viewed as narrow because, in reality, patient engagement has many iterations, setbacks, and unintended outcomes. Although the metrics are formulated neutrally to capture both positive and negative effects, we acknowledge this approach may limit the flexibility to adapt to changing needs and circumstances.

CONCLUSIONS
This study describes a tailored framework that could serve as a starting point for monitoring and evaluating patient engagement initiatives. We recommend adapting and refining the framework as appropriate to enhance its relevance to the specific context and encourage collaborative reflection. Such tailored frameworks can support learning and demonstration of impact, ultimately enhancing patient engagement in health product research and development.

Patient-Friendly Recap
• Community advisory boards (CABs) are created by patient communities, health organizations, and industry sponsors to provide a structured way for disease-specific patient experts to interact with those involved in medical research and development.
• Monitoring and evaluating CABs can 1) show whether their contributions reflect the needs of the broader patient community, 2) help both industry and patients achieve their goals, and 3) identify areas where challenges remain.
• To be meaningful for those involved, CAB evaluation methods should be jointly created. Authors of this study identified a core set of data metrics, supplemented by a “menu” of flexible optional items, to monitor progress and evaluate specific impacts of individual CABs.

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Author Contributions
Study design: Fruytier, Vat, Camp, Houýez, Pittens, Schuitmaker-Warnaar. Data acquisition or analysis: all authors. Manuscript drafting: Fruytier, Vat, Camp, Schuitmaker-Warnaar. Critical revision: all authors.

Conflicts of Interest
Elena Zhuravleva is a shareholder in F. Hoffmann-La Roche Ltd.

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