'We are data rich but information poor': how do patient-reported measures stimulate patient involvement in quality improvement interventions in Swedish hospital departments?

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ABSTRACT
Objective This study aimed to investigate if and how patient-reported measures from national and local monitoring stimulate patient involvement in hospital quality improvement (QI) interventions. We were also interested in the factors that influence the level and degree of patient involvement in the QI interventions.

Methods The study used a qualitative, descriptive design. Inspired by the Framework Method, we created a working analytical framework. Four hospital departments participated in the data collection. Collaborating with a QI leader from each department, we identified the monitoring systems for the patient-reported measures that were used to initiate or evaluate QI interventions. Thereafter, the level and degree of patient involvement and the factors that influenced this involvement were analysed for all QI interventions. Data were mapped in an Excel spreadsheet to analyse connections and differences.

Results Departments used patient-reported measures from both national and local monitoring systems to initiate or evaluate their QI interventions. Thirty-one QI interventions were identified and analysed. These interventions were mainly conducted at the direct care and organisational levels. By participating in questionnaires, patients were involved to the degree of consultation. Patients were not involved to the degree of partnership and shared leadership for the identified QI interventions.

Conclusions Overall, hospital departments have limited knowledge regarding patient-reported measures and how they are best applied in QI interventions and how they support improvements. Applying patient-reported measures to hospital QI interventions does not enhance patient involvement beyond the degree of consultation.

INTRODUCTION
Healthcare organisations have increased discussions regarding the evaluation and monitoring of healthcare performance from patients’ perspectives1–3 and the transformation of these data into quality improvement (QI)4,5 and safety improvement interventions.1,6 It is well established that healthcare is experiencing challenging times in balancing the population’s increasing and changing demands with limited finances for digital and scientific development. During the past decades, healthcare objectives have transformed from mostly managing patients with acute injuries or illnesses to providing long-term support of patients with multiple chronic conditions and managing public health at the governance level. Consequently, healthcare outcomes are no longer clearly demarcated, and they extend beyond medical goals.7 Thus, current approaches to QI constantly need revision.8

Therefore, patients and the public are key contributors to identifying areas of QI that meet patient priorities and improve
Despite the lack of consensus on the definitions of patient and public involvement, it is an important component that makes a difference in high-quality healthcare. In direct care, patient priorities combine individuals’ specific realistic health goals based on what matters most with the healthcare activities they are willing and able to perform to achieve their goals. However, patients are involved in their own care and at multiple levels of healthcare; in direct care, organisational design and governance, and policy-making. QI interventions must be executed at multiple levels of healthcare, and therefore, different collaborative approaches to patient involvement in QI, such as co-design and co-production of healthcare services, are being explored.

Discussion regarding the importance of relevant monitoring, control and agreement of the proper standards and measurable indicators has followed the development towards a more participative era within healthcare. Simple and crude patient satisfaction scales have advanced to become more patient centred, including the use of patient-reported experience measures (PREMs) and patient-reported outcome measures (PROMs). Patient-reported measures collect health outcomes directly from the patients and the public who experience them to support clinical decision-making and prioritisation of patients, compare outcomes of healthcare providers, stimulate QI and evaluate practices and policies. However, this strategy is not without opposition, and it has been argued that measurements risk becoming too excessive, standardised or generalised to identify QI areas from patient and population perspectives. Furthermore, there is little knowledge regarding the distinctions and associations between PREMs and PROMs and their actual effect on patient outcomes. Thus, reliable and valid tools for measuring patient and public involvement are also being investigated.

In Sweden, the healthcare system is tax financed, decentralised and politically steered. The health and welfare system finances and provides almost all healthcare services covering all residents, with costs heavily subsidised and no additional private healthcare insurances required. The role of the state is to legislate and establish principles or guidelines, distribute responsibilities, allocate government grants and supervise and decide on local government financial equalisation and high-cost protection/maximum fees. Since 2009, the National Patient Survey (NPS) has consistently collected PREMs at the national level in Sweden. The purpose of the NPS monitoring is to initiate QI only and enable healthcare development from the patient perspective, to facilitate evaluations of healthcare settings and provide tools for quality management. County councils, regions and local healthcare units also conduct ‘in-between measurements’, which are additional patient-reported measurements in their own contexts. Furthermore, national quality registries (NQRs) continuously report PREMs and PROMs. In Sweden, there are more than 100 NQRs. An NQR contains structured, individualised medical data for a specific patient population or for patients undergoing a specific healthcare process. Healthcare providers routinely collect data to initiate and monitor QI. For an NQR to be certified at a high level, PROMs are obligatory. To capture the patients’ perspectives, it is recommended that an organisation of patient representatives and professionals support each registry and share joint responsibility for its development. Approximately 90% of the NQRs include some form of PROMs (generic or disease/symptom specific), and about 40% include PREMs. These measures potentially impact QI at different organisational levels of healthcare.

However, there is significant opportunity for improvement. Research shows there is limited evidence regarding the ways that aggregated patient-reported measures inform QI interventions in practice. Some barriers exist, such as if and how patient-reported measures actually contribute to patients’ active choices and the perceived scepticism of professionals. In addition, there lacks explicability and timeliness of the patient-reported measures as well as structure, support and guidance in the transformation process. Research also highlights problems with identifying the monitoring systems that can be applied in practice and the patient-reported measures that provide the relevant input for each QI context when organising and managing patient involvement in hospital QI interventions. Thus, the objectives of this study were to identify the national and local monitoring systems containing patient-reported measures that are available in hospital departments and to investigate how they are applied to QI. Furthermore, we sought to explore how patient-reported measures stimulate patient involvement in QI interventions in practice. We used the following research questions:

- What monitoring systems of patient-reported measures are used?
- What QI interventions have been initiated or evaluated in applying patient-reported measures?
- How do patient-reported measures stimulate patient involvement in hospital QI interventions?

**METHODS**

**Patient and public involvement**

This qualitative study has an explorative, descriptive design. The study is part of and informed by a larger research project to study patient and public involvement in the QI interventions of hospital organisations from the clinical microsystem and leadership perspectives. Patients were not involved in the design, recruitment to or conduct of this study. However, patients participated in the larger research project. Results from this study will be disseminated to participants of the research project on publication.

**Participating settings and departments**

Two mid-size, non-academic hospitals in two different regions in southern Sweden (hospital 1 and hospital 2)
were initially considered for the study. Hospital 1 provides healthcare in all specialties to approximately 300,000 citizens in the region. At the time of the study, the hospital applied the Hoshin Kanri35 approach to its strategic planning and follow-up. Organisations that use Hoshin Kanri often follow a plan comparable to Deming’s Plan-Do-Study-Act cycle.36 The hospital applied an X-matrix to ensure that all the organisational levels effectively visualised management’s three focus areas, of which one was phrased, ‘Together with the patient—for the patient’. Hospital 2 provides healthcare in all specialties to approximately 365,000 citizens. Aligning with the region in general, the hospital applied the balanced scorecard37 to its performance management, which provides strategic financial and non-financial performance measures that enable the hospital to better accomplish its objectives. The balanced scorecard defined five perspectives, of which one was phrased ‘The citizen and customer perspective’.

Procedure

Using the Framework Method38 as a starting point, two authors (CB and CP) created a working analysis framework (table 1) adapted from the ‘Multidimensional Framework for Patient and Family Engagement in Health and Health Care’ framework.15 Stages 4–738 were completed to develop and apply the analysis framework, chart the data in the framework matrix and interpret the data.

To identify departments eligible for inclusion and their local QI leaders, we contacted the development managers (n=5) for the two hospitals, informed them about the study and asked them to forward our request to their local QI leaders or provide us with their contact information. Subsequently, each department’s QI leader(s) were informed about the study and asked to participate, and the analysis framework was provided to them in advance. Seven departments agreed to participate from hospital 1 (n=2) and hospital 2 (n=5). After a closer review, three of the departments had no practical experience with applying patient-reported measures to their local QI interventions, and consequently, they were excluded from participation. Eventually, four departments (and their QI leaders) from hospital 2 participated in the study. The participating departments represented internal medicine, oncology, paediatric and rehabilitation. All QI leaders had experience with QI interventions that were initiated or evaluated with patient-reported measures monitored at different levels in their departments.

Before commencing the data collection, the analysis framework was tested with the QI leader from the first department to participate. Specifically, a QI intervention with no degree of patient involvement on the engagement continuum was identified, and the analysis framework was adjusted accordingly. No further adjustments were made. Thereafter, one author (CB) collected data in collaboration with the QI leaders from the other three departments. These data collection meetings occurred at the QI leaders’ worksites and were digitally recorded in July and August 2021. In these meetings, we identified the monitoring systems containing patient-reported measures used. Next, we analysed the information to determine if the patient-reported measures had initiated or evaluated any QI interventions. Each QI intervention was analysed to determine the level and degree of patient involvement on the engagement continuum. Finally, we discussed the factors that influenced patient involvement.

| Factors influencing involvement | The factors that influence patient involvement in the local QI intervention. | What factors may have influenced the adaptation of patient involvement in the local QI intervention? |
|---------------------------------|---------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|
| Dimensions                      | Definition                                                                 | Analysis questions                                                                                   |
| Level of involvement            | The level of the healthcare system at which the QI intervention is carried out. | At which level of the healthcare system is the QI intervention carried out? 1. Direct care (eg, individual patient care). 2. Organisational design and governance (eg, QI team, patient safety team, hospital-wide project, leadership and management). 3. Society (eg, national health programme, policy-making, research). |
| Degree of involvement           | The place on the continuum of patient involvement where the QI intervention is carried out. | Where on the continuum of patient involvement is the QI intervention carried out? No patient involvement. 1. Consultation (eg, informed, survey, focus group). 2. Involvement (eg, invited to give feedback, active involvement, patient representative, QI team member, steering groups, research). 3. Partnership and shared leadership (codesign and coproduction, coleadership in QI). |

QI, quality improvement.

Table 1 The working analysis framework adapted from the ‘Multidimensional Framework for Patient and Family Engagement in Health and Health Care’15
Factors that influence patient involvement

In the data collection meetings with QI leaders, the factors they believed influenced patient involvement in the local QI interventions were discussed from the perspective of the patient, the organisation and society. Patients were consulted regarding outcomes of NQR monitoring, NPS and local patient surveys. Unlike the other departments, paediatric patients (ie, their family members) were actively involved in QI interventions, such as a breastfeeding project in the neonatal intensive care unit and a project to improve at-home paediatric healthcare. Patients were not involved in QI interventions to the degree of partnership and shared leadership in any of the four departments.

RESULTS

Monitoring systems with PREMs and PROMs and their connections to QI interventions

The four departments applied patient-reported measures from nine different national monitoring systems (seven NQRs, one Patient Care Bundles monitoring and one from the NPS) and several local measures and patient surveys to initiate or evaluate their QI interventions (table 2). The departments applied national and local monitoring to the same extent, but the oncology department used local monitoring more frequently. A total of 31 (range 5–11) QI interventions were identified (table 2, table 3), three of which applied a combination of national and local monitoring.

The analysis of the ratings for each QI intervention (table 4) indicated that the internal medicine department performed all QI interventions at the level of direct care. To initiate or evaluate QI interventions, outcomes from the national guidelines for the standardisation of care paths (Patient Care Bundles) and local patient surveys were applied. Thus, patients were consultatively involved in QI interventions. The rehabilitation department performed QI interventions at the direct care level and the organisational level. Patients were mainly involved to the degree of consultation and represented by the outcomes of NQR monitoring, but in one QI intervention, patients were not involved at all. The oncology department conducted QI interventions at all hospital levels but mainly at the organisational level. Patients were consultatively involved in QI interventions from the outcomes of NPS and local patient surveys. However, patients were not represented in any degree in two cases. The paediatric department performed QI interventions at the organisational and policy-making levels.

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systems to retrieve information, initiate QI interventions and enable learning. National and local monitoring were occasionally inconsistent. Patient-reported measures at the national level were excessively aggregated, and consequently, professionals questioned them as indicators for local QI interventions and suggested that the measures

| Monitoring system | QI interventions initiated or evaluated |
|-------------------|----------------------------------------|
| National Quality Registers (n=7): | Improving the pain rehabilitation process. |
| ► The National Quality Registry for Pain Rehabilitation (NRS). | Improving patient satisfaction with the rehabilitation programme. |
| ► The WebRehab. | Increasing involvement of next of kin in the rehabilitation programme. |
| ► The Swedish National Diabetes Register (NDR). | Mapping the clinical microsystem as part of a reorganisation (rehabilitation). |
| ► The Swedish Rheumatology Quality Register (SRQ). | Diabetes project—improving diabetes care. |
| ► The Swedish Web System for Enhancement and Development of Evidence-based Care in Heart Disease Evaluated According to Recommended Therapies (SWEDEHEART). | Development of register questionnaire (rheumatology). |
| ► The Neonatal Quality Register (SNQ). | Improved, person-centred medical treatment process (heart diseases). |
| ► The Swedish National Airway Register (SNAR). | Breastfeeding project (neonatal intensive care unit). |
| ► Improving the pain rehabilitation process. | Improving cooperation of health and welfare providers (asthma, COPD). |
| ► Increasing patient satisfaction with the rehabilitation programme. | |
| ► Increasing involvement of next of kin in the rehabilitation programme. | |
| ► Mapping the clinical microsystem as part of a reorganisation (rehabilitation). | |
| ► Diabetes project—improving diabetes care. | |
| ► Development of register questionnaire (rheumatology). | |
| ► Improved, person-centred medical treatment process (heart diseases). | |
| ► Breastfeeding project (neonatal intensive care unit). | |
| ► Improving cooperation of health and welfare providers (asthma, COPD). | |

| National Patient Survey | Improving the information to patients about medical students in the department. |
|► Improving patient-centredness by implementing standardised questions of ‘what matters to you’. | Improving patient-centredness by implementing standardised questions of ‘what matters to you’. |
|► Evaluation of the local ‘My care plan’. | Improving the care for children with failure to thrive. |
|► Improving healthcare at home. | Improving healthcare at home. |

| Patient Care Bundles—lung cancer | Increasing patient involvement and shared decision-making. |
|► Improving outcomes of health self-assessment. | Improving the emotional support. |

| Local monitoring of operations | Benchmarking to decrease variation between local practices’ goal formulation with patients. |
|► Improvement project concerning integrated care for children with complex needs. | |

| Local monitoring—direct patient care | Improving the identification of what matters to each individual patient. |

| Local monitoring—self-assessed health | Improving the identification of individual care needs of each patient. |
|► Implementation of a ‘Fatigue school’. | Improving outcomes of health self-assessment. |
|► Development of a programme for physical activity and training. | Improving the emotional support. |
|► Development of an information process to patients about physical activity and training. | |

| Local patient survey | Radiation therapy ‘drop-in’ project. |
|► Improving the patient’s physical path in the radiation treatment reception. | Improving the emotional support. |
|► Evaluation of local ‘My care plan’. | Development of a programme for physical activity and training. |
|► Person-centred care project: improving patient admission routines by coordinating doctors and nurses. | Development of an information process to patients about physical activity and training. |
|► Implementation of an introductory conversation about the radiation therapy. | Improving the emotional support. |
|► Children with failure to thrive—improving care. | |
|► Improving healthcare at home. | Improving healthcare at home. |
|► Improving discharge routines. | Improving discharge routines. |

| Local patient surveys on the national website “1177” for public healthcare guidance | Improving the use of the online patient dossier platform for communication between patients and professionals. |

COPD, chronic obstructive pulmonary disease; QI, quality improvement.
be used only for benchmarking. Therefore, local patient surveys were preferred to evaluate QI interventions. QI leaders suggested that departments focus more on the complex PREMs (satisfaction scales) than the PROMs, and do not consider the use of a combination of the two. Furthermore, it was difficult to connect local QI interventions to improved patient-reported outcomes and obtain reliable evidence bases. In most QI interventions patients were not actively involved because their preferences were believed already identified from the registries and surveys. Overall, departments had limited knowledge on how to apply the various patient-reported measures and how to actively involve patients in QI interventions.

From the perspective of society, which includes social norms, regulations and policy, the accreditation systems were highlighted as important factors that controlled which patient-reported measures the departments addressed and which QI interventions the departments initiated. Furthermore, the QI leaders considered the scientific evidence and monitoring grounded in nationally established policies and programmes as significant factors. However, the national goal to standardise patient processes, measurements and monitoring to simplify and make them more homogeneous could, according to the QI leaders, potentially undermine other healthcare goals, such as equitable care for citizens and more participative, co-produced healthcare.

**DISCUSSION**

**Hospital departments are data rich but information poor**

This study aimed to investigate how patient-reported measures from national or local monitoring stimulate patient involvement in hospital QI interventions. We were also interested in the factors that influence patient involvement in the QI interventions. The data collection reflected that hospital departments are data rich but information poor because they have a significant amount of data available from national and local monitoring systems yet limited resources (time, knowledge and motivation) to transform the data into QI practice and knowledge. As previously mentioned, there are more than a hundred NQRs in the Swedish healthcare context.\(^\text{28}\) Approximately 90% of the NQRs include PROMs, and about 40% include PREMs to which NPS\(^\text{27}\) and innumerable local measurements can be added. These measures potentially influence QI interventions, but this study corroborates previous research\(^\text{30}\) and illustrates significant opportunity for improvement. Earlier research indicates that hospital professionals do not request large amounts of data as they need only the relevant data for guidance, recommendations and prioritisation of each case.\(^\text{40}\) Patient-reported measures must help professionals focus on what matters most in QI rather than overwhelm them with information or demanding administration. Professionals need to be informed of the actions that leverage their time and attention and increase patient value.\(^\text{1}\) Without relevant monitoring, the feedback loop to inform and learn from QI interventions and evaluate the effectiveness of the outcomes is missing.\(^\text{41}\)

Similarly, other research also shows that monitoring must be relevant from the patient perspective. Patients are unwilling to provide large amounts of data if they do not understand the purpose.\(^\text{40}\) Patients are more interested in whether the data reflect their needs, and they are concerned that their personal data may be neglected if only professionals define, choose or prioritise the data.\(^\text{40}\) Furthermore, patients feel they are excluded from contributing their knowledge in QI interventions because they lack professional, technical or organisational understanding.\(^\text{10,40}\)

**The nebulous connection between PREMs and PROMs and hospital QI interventions**

Conducting this research meant exploring a nebulous, complicated area with no linear connections between patient-reported measures and QI interventions. When applied, departments used patient-reported measures to initiate or evaluate QI interventions or to do both in

| Hospital department | Monitoring system and number of QI interventions | Total number of QI interventions (n=31) |
|---------------------|-------------------------------------------------|--------------------------------------|
| Internal medicine   | Patient Care Bundles (n=3) Local monitoring (n=4)| n=7                                  |
| Oncology            | National Patient Survey (n=2) Local monitoring (n=8)| n=11                                 |
| Paediatric          | National Quality Registers (n=5) Local monitoring (n=1)| n=8                                  |
| Rehabilitation      | National Quality Registers (n=4) Local monitoring (n=1)| n=5                                  |

QI, quality improvement.
| Hospital department | QI intervention                                                                 | Level of patient involvement* | Degree of patient involvement† |
|---------------------|---------------------------------------------------------------------------------|------------------------------|--------------------------------|
| Internal medicine   | Increasing patient involvement and shared decision-making                        | 1                            | 1                              |
|                     | Improving outcomes of health self-assessment                                     | 1                            | 1                              |
|                     | Improving the emotional support                                                  | 1                            | 1                              |
|                     | Improving the identification of what matters to each individual patient          | 1                            | 1                              |
|                     | Improving the identification of individual care needs of each patient            | 1                            | 1                              |
|                     | Improving discharge routines                                                     | 1                            | 1                              |
|                     | Improving the use of the online patient Dossier platform for communication between patients and professionals | 1    | 1                              |
| Rehabilitation      | Improving the pain rehabilitation process                                        | 1                            | 1                              |
|                     | Increasing patient satisfaction with the rehabilitation programme                | 1                            | 1                              |
|                     | Increasing involvement of next of kin in the rehabilitation programme           | 1                             | No information                 |
|                     | Benchmarking to decrease variation between practices in goal formulation with patients | 2              | 0                              |
|                     | Mapping the clinical microsystem as part of a reorganisation                    | 2                            | 1                              |
| Oncology            | Development of a programme for physical activity and training                    | 1                            | 1                              |
|                     | Improving the patient's physical path in the radiation treatment practice        | 1                            | 1                              |
|                     | Implementation of a ‘Fatigue school’                                            | 2                            | 0                              |
|                     | Development of an information process to patients about physical activity and training | 2    | 1                              |
|                     | Improving the information to patients about medical students in the clinic       | 2                            | 1                              |
|                     | Improving patient-centredness by implementing standardised questions of ‘what matters to you’ | 2              | 1                              |
|                     | Radiation therapy ‘drop-in’ project                                              | 2                            | 1                              |
|                     | Person-centred care project: improving patient admission routines by coordinating doctors and nurses | 2              | 1                              |
|                     | Person-centred care project: improving information of appointed contact nurse   | 2                            | 1                              |
|                     | Implementation of an introductory conversation about the radiation therapy       | 2                            | 1                              |
|                     | Evaluation of local ‘My care plan’                                              | 3                            | 0                              |
| Paediatric          | Improved, person-centred medical treatment process (heart diseases)             | 2                            | 1                              |
|                     | Improvement project concerning integrated care for children with complex needs  | 2                            | 1                              |
|                     | Improving cooperation of health and welfare providers (asthma, COPD)             | 2                            | 1                              |
|                     | Breastfeeding project in the NICU                                               | 2                            | 1, 2                           |
|                     | Children with failure to thrive—improving care                                  | 2                            | 2                              |
|                     | Diabetes project—improving diabetes care                                         | 2, 3                         | 2                              |
|                     | Improving healthcare at home                                                     | 3                            | 2                              |
|                     | Development of the NQR questionnaire                                             | 3                            | 2                              |

*1. Direct care. 2. Organisational design and governance. 3. Society.  
†0. No patient involvement. 1. Consultation. 2. Involvement. 3. Partnership and shared leadership.  
COPD, chronic obstructive pulmonary disease; NICU, neonatal intensive care unit; NQR, national quality registry; QI, quality improvement.
some situations. This study did not map the total number of patient-reported measures available to each department. Thus, it is impossible to discuss compliance with all existing monitoring. Furthermore, we did not distinguish between the PROMs and PREMs applied. However, the results illustrate some patterns.

Hospital care for adults used QI interventions at the direct care and organisational levels, while paediatric care used QI interventions at the organisational and societal levels of healthcare (table 4). For example, the internal medicine department consistently performed QI interventions at the level of direct care and involved patients by using the survey outcomes. The oncology department predominantly used local monitoring (table 3). In the paediatric department, patients (or their family members) were consistently more actively involved. However, using the analysis framework,15 the examples of active involvement in the paediatric department were not different from other QI interventions mapped to the second degree of involvement (table 4). No direct conclusions can be drawn from this result. However, different departments had various levels and degrees of patient involvement in QI interventions, and it would be interesting to analyse these differences through additional research.

None of the departments involved patients to the degree of partnership and shared leadership15–17 in their QI interventions (table 4). Although this outcome is disappointing, it is not surprising. Swedish hospitals rarely involve patients in co-design and co-production activities16 17 with QI interventions.

This study speculated that a connection exists between the departments’ application of patient-reported measures to QI interventions and higher degrees of patient involvement. However, the results did not indicate any such connections. Rather, the analysis indicated that the QI intervention already considers the patients’ preferences by using information from the monitoring systems. Although it may be a good starting point to incorporate PREMs and PROMs from the systems, many QI interventions have local objectives. Thus, professionals should also consider the degree of patient involvement that supports the purpose of each QI intervention.18 Merely using reported data from systems does not constitute patient involvement.

Both QI leaders and development managers highlighted the low response rate from patients on local surveys as one of the influencing factors related to the patient perspective. They believed patients did not understand their own importance of giving feedback to improve the services. Similar to previous research, the leaders and managers argued that the patient’s limited feedback affects the motivation of the professionals to incorporate patient involvement in QI interventions.19 The lack of knowledge about how patients view the patient-reported indicators and how relevant the indicators are to their concerns justifies this position. Furthermore, the patient’s role in various QI interventions (and in various healthcare contexts) needs clarification.10 18 40 42 Thus, the results of this study highlight the importance studying active patient involvement in QI more intensely, and in such research, patients should be actively included.

Moreover, major influencing factors relate to the organisational and societal levels of healthcare. National regulations, programmes and accreditations significantly impact the prioritising of QI interventions.32 Obviously, a gap exists in understanding the role of the patient and the public in QI interventions, monitoring and follow-up of outcomes at all levels.10 13 32 40 Even though patient involvement in QI interventions may be a complex activity in practice, knowledge regarding QI is necessary and requires supervision and management.32 This ‘black box’ must be opened if healthcare is to become more participative and co-produced, equitable and suitable for the public’s purposes.14 16 To establish the input data relevant for each QI intervention, whether patient reported or not, further research and knowledge regarding the inter-relationships between PREMs and PROMs and the ways in which they constitute relevant input and feedback for QI interventions must be prioritised.23 43 44 Are PREMs and PROMs more suitable for monitoring at the group level and PROMs more suitable for guiding individual treatments? Are PREMs and PROMs appropriate incentives and tools to enhance patient involvement in QI interventions or are other approaches more appropriate?3–5 16 17 Is it helpful to measure patient involvement in QI interventions, and if so, how?23 These questions need further study.

**Strengths and limitations**

This study provides valuable input regarding patient involvement in QI interventions. We experienced some challenges in applying the working analysis framework to organise and manage data (table 1) in the complex context of hospital QI interventions. However, we focused on the aims of the research questions throughout the data collection and analysis. To minimise misunderstanding during the data collection, individual meetings were held with the department’s QI leaders and digitally recorded. Using the Framework Method38 provided a systematic and flexible structure to manage and guide the data analysis and reporting and counteracted the challenges.

Some of the issues in conducting the research need elaboration. It became apparent at the beginning of the study that hospital departments generally had limited knowledge regarding patient-reported measures and the concepts of PREM and PROM. We approached QI leaders after contacting the development managers, but despite reminders, only seven departments responded positively. Moreover, only four of the responding departments (all from hospital 2) had practical experience applying patient-reported measures in their QI interventions. At the time for the study, hospital 1 implemented a major organisational change that may have affected its response rate. Furthermore, we contacted the leaders during the summertime and the ongoing COVID-19 pandemic.
However, we were not convinced that these circumstances explained the significant lack of responsiveness.

Therefore, we contacted the development managers again, and their responses confirmed our presumptions that attributing the lack of responsiveness to the ongoing COVID-19 pandemic was a simplified explanation. Instead, the general opinion was that hospitals are consistently under significant pressure, and QI leaders do not have time to routinely reflect on overarching, strategic questions. Furthermore, QI leaders are not educated to systematically apply and learn from healthcare outcome monitoring and QI interventions. The development managers also indicated that using the patient involvement approach for QI further complicated the study assignment for QI leaders because patient involvement is neither well defined nor fully applied in hospital organisations. The feedback from the development managers corroborates earlier research that indicates current measurements and monitoring are problematic to apply and enact in complex healthcare settings. Consequently, hospital organisations do not maximise their potential for QI. Patient processes, measurements and monitoring are being standardised to address these issues, but this standardisation adds a risk of information being oversimplified and professionals underusing it. QI interventions and (patient involvement in QI interventions) are afterthoughts to the daily work and knowledge, learning and improvement in this area stay limited.

CONCLUSIONS

The results from this study indicate that hospital departments generally have limited knowledge regarding patient-reported measures, how best to apply them in QI interventions and how the measures contribute to improvements. If applied at all, patient-reported measures are mainly used in QI interventions performed at the direct care and organisational levels. Patients are involved to the degree of consultation mainly by participating in surveys. However, paediatrics provides examples of patients (or family members) being actively involved in QI interventions that patient-reported measures initiated or evaluated. National patient-reported measures and local patient surveys do not yet encourage patient involvement to the degree of partnership and shared leadership in hospital QI interventions.

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Patient consent for publication Not applicable.

Ethics approval The study is part of a larger research project at the University of Borås and Jönköping University. Because individuals were involved in the data collection of this study, efforts were made to carry it out according to current laws, regulations and codes of research ethics. Therefore, and prior to commencing the research project, permission was obtained from the Regional Ethical Review Board in Gothenburg, Sweden (Dnr: 1006-16) with a complementary permission from this study (Dnr: 2021-01456). Informed, written consent was obtained from the participants.

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