Challenges and facilitators for patient and public involvement in England; focus groups with senior nurses

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ABSTRACT

The concepts of patient and public involvement (PPI) have been recognized and linked with quality in health services internationally and in Europe. In England, for more than a decade, NHS policies have increasingly quoted patient-centred services. Limited evidence exists about the implementation of PPI policies and strategies within organisations; three studies only have explored health professionals’ perceptions of PPI. Although nurses’ positive support for patient and public involvement has been noted, comparatively little is known about senior nurses’ experiences of embedding PPI. A national consultation utilising three focus groups aimed to explore senior nurses’ perceptions of challenges and facilitators for PPI implementation. Four Strategic Health Authorities (SHAs) and eleven Primary Care Trusts (PCTs) in England, with fifteen senior nurses with leadership roles and direct PPI experience, participated. Nurses’ perceptions on patient and public involvement, challenges and facilitators for its implementation were discussed. Focus groups were digitally recorded and transcribed verbatim; anonymised transcripts were validated by participants and analysed with thematic analysis. Limited resources, patient representation and recruitment, complexities of implementing PPI and national policy changes were challenging. Commissioning limitations, lack of feedback on patient experience, limited staff awareness, negative attitudes, management of patients and public expectations constituted further challenges. Nursing role characteristics and informal involvement activities, PPI policy and cultural change, commissioning PPI competencies, related service frameworks, providing feedback on patient experiences to staff and recognition of involvement benefits were recognised as facilitators. Findings provided new insights into senior nurses’ experiences and evidence that progress towards meaningful, effective PPI remains slow. However, recognition of existing nursing role characteristics and potential delivery problems created by expanded nursing roles, informal PPI practice and internal organisational sharing of patient feedback may bring an “emerging productive partnership” with nurses enabling and contributing to effective PPI.

Keywords: Nursing; Patient and Public Involvement; Challenges and Facilitators; Focus Groups; Patient-Centred Care

1. INTRODUCTION

The concepts of patient and public involvement (PPI) and empowerment have been recognized and linked with quality in health services internationally and in Europe [1,2]. Countries have implemented a wide range of patient empowerment measures, including patients’ rights legislation (Netherlands, Greece), introducing ombuds-person services (Austria, Finland, Hungary, Norway, Greece) and increasing patients’ involvement and participation in care decision-making (England) [3,4].

In England, for more than a decade, NHS policies have increasingly quoted patient-centred services. Notable drivers have been the legal duty to involve and consult the public [5] and the increasing body of international evidence for involving people in health care and its benefits [6-8]. Most recently, the PPI agenda has permeated the World Class Commissioning vision, stating that “to be world class commissioners we need to know the needs and preferences of our local communities, work with our partners on the health and well-being agenda and work with local people to tackle health inequalities”. Specific emphasis was placed on “building continuous and meaningful engagement with
patient and public to shape services”, as one of its competencies [9]. Furthermore, the “High Quality Care for All” [10] called for an NHS “that gives patients and the public more information and choice, works in partnership and has quality of care at its heart”.

Many definitions exist for involvement; INVOLVE [11] summarised this as “everything that enables people to influence the decisions and get involved in the actions that affect their lives”. To enable policy implementation, the English NHS has adopted the “Involvement Continuum” [12], encompassing giving and getting information, forums for debate and participation (Figure 1).

Although the research evidence supporting PPI varies in quality, benefits identified in systematic reviews have included improvements in health literacy, clinical decision making, self-care, chronic disease management and patient safety [13,14]. Others have also demonstrated improvements in the information clarity provided and patients’ knowledge [15]. However, limited evidence exists about the implementation of PPI policies and strategies within organisations [16]. National surveys found that although health services were putting systems in place to involve people in planning and improving services, many challenges remained. Barriers included lack of commitment by senior managers or clinicians, inadequate resources, lack of incentives and poor quality of interpersonal care [17-20]. Other studies have also found that health services staff were often the most important change drivers, but strong management and leadership, acting on patient feedback and strengthening an open and transparent culture were also vital [21].

Only three studies have explored health professionals’ perceptions of PPI [22-24] showing that progress towards achieving meaningful and effective PPI was slow. Deficiencies in financial and human resources, organisational capacity, lack of relevant data, difficulties in supporting the public and accessing seldom heard groups were identified as barriers. Significant changes to the way that PCTs organised PPI in commissioning, indicated the start of a cultural shift; however, engaging beyond the ‘easy to reach’ was cited as a barrier [23,24]. An ethnographic study [23] found that health professionals determined areas for service user participation, which covered a wide range of activities; understanding and practice relating to this varied according to professional ideologies and circumstances. An interesting finding was the challenge embedding user involvement across work streams and authority issues raised.

Comparatively little is known about senior nurses’ experiences of embedding PPI, although nurses’ positive PPI support has been noted [6]. Nurses are key NHS frontline staff in terms of direct patient care; they also

![Figure 1](https://via.placeholder.com/150)

**Figure 1.** The involvement continuum.

hold key management positions which require an understanding of PPI and its policy implementation challenges. This article reports on a national consultation exercise commissioned by the Royal College of Nursing (RCN) to explore senior nurses’ perceptions of PPI challenges and facilitators in England. Further details of the project can be found in the full report [25].

### 2. METHODS

A qualitative exploratory design utilising focus groups with senior nurses—nurses with leadership roles—across England, was employed. Focus groups have the advantage of making use of group dynamics to stimulate discussion, gain insights and generate ideas to pursue a topic in greater depth. It is a useful technique for exploring values and beliefs about health, disease and systems; they are popular in health promotion and action research, organisational research and development [26]. Being relatively economic and gathering views of many in a relatively short space of time were additional reasons for choosing focus groups [27].

SHAs in England were selected as the first points of contact, having the advantage of being able to access all PCTs and a range of senior nurses across England. All nine SHAs were invited to send representatives to participate in focus groups. Letters informing and inviting senior nurse managers were sent directly from the RCN to Chief Executives for approval; they allocated a key stakeholder to co-ordinate and explore the feasibility of a focus group locally. Stakeholders invited senior nurse managers to participate voluntarily. Recruitment was very slow, although the Chief Executives and key stake-
holders were reminded with second letters and follow-up phone calls; reasons given were work overload and other priorities. All communication regarding recruitment and selection took place between June 2008 and October 2008. Ethical approval from a Trust or an external body was not required; the University Ethics Committee was informed for this national consultation.

All participants were given an information sheet and consented in writing to participation and digital recording of the focus group. Confidentiality was ensured and all participants were aware of their right to withdraw. A semi-structured topic guide was drafted in collaboration with RCN and used for all focus groups; they were facilitated by MB, a university researcher, independent from the RCN, SHAs and PCTs. They lasted between 80 and 120 minutes and took place in private meeting rooms at SHAs’ premises.

Digital recordings were transcribed verbatim by professional transcribers. Data was analysed manually by MB using the principles of thematic analysis, thus coding data line by line and forming codes, categories and themes [28,29]. Fifteen sub-themes emerged within the main themes comprising challenges (9 sub-themes) and facilitators (6 sub-themes).

Transcripts were checked against recordings for reliability. Anonymised summaries of the individual focus groups findings were sent to participants of each group for participant validation. Only two participants responded citing no comments; one of them reinforced a suggestion already made at the group.

3. RESULTS

Fifteen senior nurses, with leadership roles and direct PPI experience, employed in eleven PCTs from four SHAs participated. Participants, all RCN members with a variety of roles (Table 1), took part in three focus groups at north, west central and south England locations, conducted between September and October 2008.

3.1. Challenges for Effective PPI Nursing Practice (Figure 2)

3.1.1. Limited Resources, Capacity and Time

The importance of resources was highlighted among all groups. On one hand, the organisational rationing of general resources allocated to specific projects and events including PPI, was discussed. On the other, allocated PPI resources, capacity and time for PPI activities were also recognised as limited. These limitations also affected the time for change implementation emerging from PPI; in one case, it had taken three to four years for the recommendations to be implemented. Additional PPI capacity for specific services and subgroups of service users, i.e. children, was also needed.

Figure 2. Challenges for patient involvement.

Table 1. Professional roles of participants.

<table>
<thead>
<tr>
<th>Focus Group 1</th>
<th>Focus Group 2</th>
<th>Focus Group 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Director of Quality and Performance (Executive Nurse)</td>
<td>Clinical Audit Manager</td>
<td>Lead Infection Policy Chief Nurse</td>
</tr>
<tr>
<td>Head of Service Development (Children Services)</td>
<td>Lead Educator</td>
<td>Director of Nursing (Executive Nurse)</td>
</tr>
<tr>
<td>Head of Clinical Strategy</td>
<td>Director of Nursing</td>
<td>Head of Clinical Governance and Effectiveness</td>
</tr>
<tr>
<td>Head of Infection Prevention and Control</td>
<td>Head of Clinical Services</td>
<td>Head of Service Reform</td>
</tr>
<tr>
<td>Commissioning Manager (Sexual Health)</td>
<td>Matron (Senior Clinical Nurse)</td>
<td></td>
</tr>
<tr>
<td>Public Involvement Lead and Health Improvement Manager</td>
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Participants spoke extensively about availability of resources, capacity and time at a workforce planning level especially at the initial PPI stages and development, and also through the different levels of the involvement continuum. Resources were also important when “real” involvement and partnership had been “achieved”, for sustainability, as patients’ expectations from the services might have been increased.

Well as you work up the continuum again, if you’re working towards real involvement and real partnership, it takes time and to go out to community groups, to link with Sure Start, to set up your days, to… The maternity
services are like the Mental Health Services, they’ve got a long history of involvement, haven’t they? And it takes time to work in partnership with the public in whatever way is appropriate for that particular issue or service.

(FG2, P4, p24, 19-25)

Effective management, setting up and maintenance of memberships, i.e. of Patients Forums, and involving everyone individually also required staff time and resources. These elements should also be seen in conjunction with the other responsibilities of nurses, especially when staff shortages were considered. The additional nursing roles and absorbing different roles with various deliverables could be challenging for PPI.

I mean as nurses have taken on more and more of the medics roles, which they have but actually even with skilled nurses, they still haven’t dropped a lot of the other roles that they had… And to a certain extent I think a lot wouldn’t want to as well, it comes to a point where the NHS has worked on goodwill for years, hasn’t it? We’ve all done it but this, you start to get to a point where sometimes it’s just that bit too much and people are starting to get a bit more protective of their time.

(FG1, P3, p27, 24-34)

3.1.2. Representation and Recruitment of Patients and the Public

High representation of people with certain characteristics in conjunction with their self-selection, were discussed extensively. Patient representatives were often people who had time, were more vocal and active, and perhaps had an interest in specific issues. Other people were less represented, i.e. those who could not read English, were less educated or illiterate, less well off, employed or with childcare issues. Thus, as a result the same patients were involved in all activities creating “real” representation challenges.

I’m just thinking about some of the parents we’ve got who can barely read, who are very hard to reach and who would be completely fazed by having to go through that kind of process but probably wouldn’t go through it anyway, we already know that those who are not so well off are less likely to attend because either they’re working or they’ve got issues of, childcare issues or this, that and the other.

(FG1, P3, p27, 24-34)

Recruiting and influencing people to participate using a variety of approaches were also challenging. Lack of guidelines for getting a wide range of people was highlighted. Involving the public, “the man in the street” in particular, presented a challenge for many participants. Involving certain population groups and its organisational challenges were also discussed. For example, the difficulties and time required for children and young people’s involvement may be challenging to particular groups of professionals, i.e. children’s nurses. Additional challenges were associated with staff being equipped and trained to deal with black and ethnic minorities (BME) and the disabled, i.e. answering telephone lines appropriately and presentations with interpreters were cited.

I think a couple of challenges from my perspective, the BME community and the deaf, the hard to reach communities, because there’s nothing more terrifying than standing up doing a presentation, seeing it sign languaged, hearing it being interpreted in Cantonese and seeing somebody else, you know, and are we equipped and trained to do that?

(FG3, P1, p17, 6-10)

3.1.3. Complexity of PPI Implementation

The plethora of complex dimensions and involvement areas, i.e. many clinical practices, various professional cultures and subsequent organisational issues created additional difficulties. It was recognised that changes and development had to be implemented across the spectrum of clinical services and professions, so cultural change could be facilitated. The additional complexity of PPI being linked with people’s immediate personal experiences, sometimes with unpredictable events, was also recognised.

Set up a consultation event, here’s the ground rules, here’s the parameters, here’s what we will do with your information, here’s what we (it is easy), but we’re talking about day to day lived experiences that pop up when you least expect them, or you get put in a situation where you hear something.

(FG3, P1, p17, 6-10)

3.1.4. National PPI Policy and Changes in Structures

Additionally, concerns were expressed that changes in national PPI policies and bodies, notably, the abolishment of Community Health Councils and PPI Forums and the creation of LINKs, had temporarily brought wider engagement to a halt.

...because we had a very active Patient and Public Involvement Forum, the LINKs host wasn’t appointed until the beginning, their contracts started at the beginning of August, then the manager within the host… host organisation was identified to start the work went off sick, you know, and now we’re here into September and really, and there’s a sense of frustration building amongst particularly the old PPI forum members, so here we are six months on from the abolishment of the forum and they still haven’t got an engagement group.

(FG2, P2, p20, 44-51 and p21, 1-4)

NHS structural changes also affected the implementation of the world class commissioning competencies, as potential collaborations with patient representatives were interrupted. These changes had also led to organisational and team changes that subsequently had a PPI impact, i.e.
amalgamation or restructuring of teams impacted on PPI. Additionally, the Patient Advice and Liaison Services (PALSs) were not considered well-integrated within the whole NHS system; dealing with patients with specific issues was considered to limit PALS’ objectiveness of feedback.

3.1.5. Commissioning Limitations: Organisational Issues
It was believed that PPI projects were still very tokenistic; there were some good examples, but the world class commissioning PPI competencies, had not yet been reached. If the organisational participation culture, as revealed from the organisational strategy and actions, was not open and transparent, then PPI was not perceived to be effective. Control and representativeness were other issues to be considered. PPI controlled either by committees or people, staff or patient representatives was considered problematic; this might be linked with user representation issues discussed.

“I think the committees, it’s how the organisation is set up, you can have controlling PPI, can’t you? You can have… it was Dignity and Compassionate Care group we’ve just set up and someone suggested a name and obviously new into the organisation I didn’t know who this person was, so I did a bit of, well, you know, she’s been around a few years, she’s been awarded an OBE for her contribution to the NHS but, you know, does she truly represent a user? So, you know, it’s that control, you can have a control system that says yes we have got PPI but then when you sort of dig down, you think is that really Joe Bloggs on the street’s view?"

(FG2, P3, p21, 21-33)

3.1.6. Staff PPI Awareness and Negative Attitudes
Participants across groups recognised low PPI awareness among nurses. This was expressed with nurses performing patient involvement activities, i.e. informing, sharing and discussing care issues with patients, without recognising or labeling them as such. They actually welcomed the informal and flexibility; they were resistant in such labels.

“…some of our consultants are fabulous, absolutely fabulous and we’ve got certain dynamic groups, diabetes and various other things but we’ve got others where the patients would never be, have a voice if it was down to the doctor. Whereas I don’t think nurses think like that, so I think it is very relevant about how boards are made up, where we’re going with all of these new Trusts, who’s the decision making people and how we involve our customers, so I do think it’s quite crucial.”

(FG2, P1, p19, 40-47)

3.1.7. Lack of Feeding Back Evidence on Patients’ Experience
The mentioned resistance of nurses towards labeling certain activities as PPI had consequences on capturing and measuring patient involvement and experience properly. Thus PPI evidence was not collected; it was difficult to be identified and demonstrated, resulting to limited feedback to the management. Clinical and other decisions were based only on demonstrable or available evidence and not on anecdotal patient experiences; thus the process of shaping services accordingly was hindered.

“…actually sometimes it’s very hard to find the evidence to actually demonstrate that and say for instance there might be notice of a meeting but they might not say “xx is the patient representative”, you know, so it maybe just had a list of names… You know, how can you demonstrate this for your, this particular assessment process or that process? Actually because it’s part of what we do on a day to day.”

(FG2, P2, p14, 15-24)

3.1.8. Management of Patient and Public Feedback and Expectations
Effective management of patient and public expectations was also perceived as difficult, requiring good judgment and balancing all complexities around them. Ineffective management could cause delays in resolving issues and result to patients’ frustration.

“…and they’re still at the end of that session frustrated because we still haven’t worked out how we’re going to get them their tablets more quickly. And they’ve been really honest. I’m frustrated because I can see that it should be simple but I know it isn’t because of all these other complex things happening, therefore our partnership’s thwarted really because we can’t get to where they want to get to very easily.”

(FG1, P1, p21, 27-33)

In some occasions, utilising the feedback, expectations or outcomes from PPI activities to shape services was difficult. This was due to practical, financial or organisational reasons. Incapacity to fulfill certain expectations, because they were outside somebody’s jurisdiction was also highlighted.
P3: My concern is that we can sort them and don’t necessarily use the information we get to shape the service if we’re not careful.
P2: Or that we don’t have the resources to provide.
P3: Yes, which is where the honesty bit comes in.

(FG1, p12, 38-44)

3.2. Facilitators for Effective PPI Nursing Practice (Figure 3)

3.2.1. Nurses’ Role and Informal PPI Activities

The nurses’ skills, their informal caring and interactive activities—although not labeled as PPI were recognised as facilitators. General comments were made about nurses having transferable skills and gaining new skills all the time. The nursing caring role included continuous rapport with patients, engagement and on-going feedback from them. This feedback, including comments, compliments and complaints—when evidenced—was taken into account in shaping services.

As a profession I think it is… that caring side of nursing, that one to one rapport that you get when you’re caring for a patient and the feedback…, so I still have that engagement and I know as leaders we’re very conscious of the fact that we need to be taking on board patient stories and we need to be identifying good and bad comments, you know, that we’re getting back from the patients and acting upon them and I think that’s how we’re going to shape our services.

(FG1, P4, p2, 24-32)

Nurses were considered to conduct PPI in practice, in every single interaction with patients, even if not recognised as such. One to one and non-patronising interaction with patients, listening to communities, individual patients and working towards their best care and services were considered to be both PPI and nursing professionalism elements. They were seen to be integral to a nurse’s role at different levels and settings, i.e. in community or acute settings. However, it was recognised that this might vary in different departments, clinics, or wards. In maternity, for example, involvement might be greater than in other clinics. In addition, specific professional roles such as the modern matrons were considered having a better PPI understanding than others.

I think if you looked at the, the modern matron role that was brought in they do have an understanding of public patient involvement as well as patient safety and the governance aspects but maybe the frontline nursing staff, it’s about that one to one interaction for patients.

(FG3, P1, p27, 11-14)

3.2.2. NHS Policy Changes and Subsequent Cultural Change

It was recognised that most recent NHS national policies had been patient-centred and facilitated a cultural change, including a focus on self-care and self-management, both closely related to PPI. Although new developments, i.e. LINKs at the time, were recognised as capable of bringing changes and shaping services, lack of clarity about their processes and membership were also mentioned.

3.2.3. Commissioning PPI Competencies

The world class commissioning PPI competencies and organisational drivers for their implementation had somehow influenced and facilitated PPI. The push for integrated services and the recognition of the third sector also reinforced patient-centred services.

Well, yes, one of the world class commissioning competencies is around Patient Public Involvement… It’s about your structure as a commissioning PCT and how you’ve structured yourself and what your hierarchy looks like and how you’re going to engage the public in, right at that front end, your health needs assessment works because at the moment they’re kind of brought in after that, aren’t they?

(FG3, P1, p32, 28-36)

The organisational strategy or requirements-revealing perhaps a positive participation culture, i.e. patient representatives on the Board-facilitated higher involvement levels. Organisational structures might also enable involvement at the higher level.

I mean for instance our Board now has two patient representatives who are, and their function has evolved and their participation in that Board has evolved, so for
instance when our new PCT was first set up one of the early Board meetings, our Chairman made the comment that said this is a meeting held in public, not a public meeting.

(FG2, P2, p20, 9-13)

3.2.4. Other PPI Supportive Frameworks
Several participants referred to existing frameworks, i.e. the Essence of Care Standards, the Knowledge and Skills Framework, the Standards of Nursing, and customer care standards. Their relevant to PPI competencies or elements were identified as helpful for PPI overall. Some participants also referred to evaluation, monitoring and appraisal processes, dealing with staff developmental issues, including PPI, as PPI supportive. The consensus in one group was that the existing competency frameworks were sufficient and adding another framework would make clinicians feel overloaded. A participant felt that the continuum of involvement could be mapped through existing frameworks, although its implementation needed work.

….because you’ve got the continuum of involvement that is recognised and actually in a sense you could put your different levels of involvement as with the gradings, you could make a model basically, you could use the continuum of involvement and then you’ve got your essence of care standards, you’ve got your KSF… you’ve got your customer care standards, which is all integral into a nurses role at different levels in different ways, wherever they may be working in the community or in an acute setting or in a community hospital. So actually there is a kind of framework there already, it’s how you apply it.

(FG2, P4, p29, 25-36)

3.2.5. Recognition of PPI Benefits
Participants referred to the PPI benefits; notably that it was powerful, influential and the whole process provided the argument for cultural and organisational change. The awareness and recognition of these benefits facilitated effective PPI further.

P4: And I think nothing’s more powerful is it, than a patient or a carer challenging a consultant or a doctor or a nurse or a Social Worker?

P2: It’s very powerful.

P4: To, or the Head of Social Care, it is much more powerful and it actually speaks volumes, you know, that they will turn and pull something up…

(FG3, p4, 36-42)

It was also recognised that the core of true patient care started with asking patients’ about individual needs. Real involvement should also have a re-energising effect for everybody in the system. Other benefits included better understanding of the NHS bureaucratic system, challenging NHS staff, bringing in simple changes, speeding up procedures, making the NHS more transparent and accountable.

3.2.6 Organisational Feedback to Staff
Providing feedback to staff on various aspects of patient experiences and involvement, for example, patient experience surveys, complaints and various incidents, was considered another facilitator. Organisational feedback could be linked with action plans. The feedback provided to individual nurses could help them recognise the different aspects’ importance and any actions or resolutions necessary.

Well we do a lot of feedback back to staff on just, of things that we’ve done, so obviously our satisfaction surveys, it would, you know, it’s a key thing to go back to staff and involve them in determining an action plan. And but also things like complaints where we know we’ve made a change and on a regular basis we’ll kind of list some of those things so some of the issues that arose, so even from a complaint or a survey or an audit or somewhere that’s involved the patient and then kind of outline what was a result really, kind of what was the outcome for that?

(FG2, P5, p6, 26-33)

4. DISCUSSION
Although the consultation exercise yielded valuable insights into the evolving process of implementing PPI in NHS Trusts in England from a senior nursing perspective—which is under-researched—its limitations should be considered. Five focus groups, representative of nine SHAs, were initially intended. However, recruitment and their facilitation organised in collaboration with Trust gatekeepers, proved very difficult. Issues such as potential participants’ workload or perhaps not-treating the topic as a priority need to be considered; these may also help us understanding PPI perceptions of these professionals. The findings are based on three focus groups only, and fifteen self-selected participants representing four SHAs; this cannot preclude bias. Furthermore, participants were all in strategic senior management positions; their experiences are not necessarily representative of a wider nursing PPI experience. This consultation was conducted in late 2008. However, guidelines for PPI implementation have not been issued yet in England, and another study exploring senior nurses’ experience of PPI has not been identified internationally. Importantly, similar issues may affect implementation of patient and public involvement not only in England, but internationally.

With regard to PPI challenges, participants’ experiences were consistent with those identified elsewhere;
limited resources, time, complexities of working across service boundaries and professional cultures [22,23]. An interesting new finding was the impact on continually expanding nursing roles of taking on new PPI and other responsibilities, which could result in delivery problems. Participants also identified difficulties recruiting representative patients, with concerns expressed about hard to reach groups and the need for recruitment guidelines; these findings were consistent with those of others [22,24]. PPI control by committees and professional staff were seen to be problematic in terms of commissioning practices. Others have also found that health professionals determined the areas of participation by service users [23]. Although many medical staff were supportive of PPI, some participants had encountered unhelpful attitudes; an earlier study [6] noted similar issues raised by giving up control.

Important new findings emerged in relation to PPI facilitators. Significant facilitators identified here corresponded to some world class commissioning priorities [9]. Participants discussed services based on patient experience, people’s choices, control and more personalisation, alongside work with community partners and engagement with public and patients, all of which were highlighted within the world class commissioning vision, associated competencies in England. Similar issues have also been identified in national and international policies elsewhere [2-4]. A strong message was that another PPI implementation framework was not needed, as some PPI related benchmarks and competencies were incorporated in the Essence of Care and Knowledge and Skills Framework in England [30,31]. It appears that nurses felt overwhelmed by frameworks and regulations. A consensus existed that nurses in practice, because of the nursing characteristics and nature, performed PPI in its pragmatic sense, even if they did not recognise or label it as such. PPI, its learning and achievement has not been given sufficient recognition, nor has it been robustly shared and systematised especially within practice. This is perhaps one of the most important findings, together with that of the need for open and transparent feedback of patient experience evidence to shape services and expedite action planning at the frontline [21].

5. CONCLUSIONS

This consultation explored “an emerging productive partnership”, with regard to senior nurses enabling patient and public involvement. Notwithstanding its limitations, rich qualitative data shed light on involvement barriers and facilitators, as experienced by senior nurses in management positions, an area that is under-researched. New findings highlighted the importance of two-way feedback of evidence based on patient experience between frontline nurses and managers; the utility of existing policy and service frameworks to enable implementation of involvement mechanisms; the need for guidelines to recruit hard to reach groups and the potential for a negative impact of relevant responsibilities on expanding nursing roles. Of particular interest, was the view that nurses performed patient involvement in a pragmatic sense, by virtue of the nature of nursing.

International and national policies elsewhere [2-4], and NHS policies in England [5-10,12], place a great emphasis on patient involvement and empowerment. However, these findings suggest that although progress has been made, many challenges still remain in transforming the way that healthcare services in England are commissioned and provided, if current aspirations on shared-decision making as “the norm” are to be met [8]. Enhancing awareness about patient involvement could benefit the development of more effective monitoring and feedback mechanisms at organisational levels. In addition, the creation of nursing involvement networks may be helpful in harnessing collective experiences to benefit patient-centred services and empowerment of patients, in England and elsewhere.

6. ACKNOWLEDGEMENTS

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