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Manuscript title:
Feasibility and acceptability of the use of patient-reported outcome measures (PROMs) in the delivery of nurse-led, supportive care to women with cervical cancer

Author names:
Grigorios Kotronoulas, PhD, MSc, BSN, RN, Research Fellow in Cancer Care
Fran O'Brien, DipHE, BA (Nurs), RN, Cancer Nurse Specialist
Mhairi F Simpson, MN, BSc, RN, Nurse Consultant Cancer Care
Roma Maguire, PhD, MSc, BN, RGN, Professor of eHealth

Affiliations:
Dr Kotronoulas, (see details for corresponding author)
Ms Fran O'Brien, Wishaw General Hospital, NHS Lanarkshire, 50 Netherton St, Wishaw ML2 0DP, Lanarkshire, UK (Fran.O'Brien@lanarkshire.scot.nhs.uk); Dr Simpson, NHS Lanarkshire, Lanarkshire, UK (Mhairi.Simpson@lanarkshire.scot.nhs.uk);
Prof Maguire, School of Health Sciences, University of Surrey, Standard Buildings, 94 Hope Street, Glasgow, G2 6PH, UK (r.maguire@surrey.ac.uk);

Corresponding author:
Grigorios Kotronoulas
School of Health Sciences, Faculty of Health & Medical Sciences, University of Surrey
Standard Buildings, 94 Hope Street, Glasgow, G2 6PH
T: +44 (0) 141 249 0922
E: g.kotronoulas@surrey.ac.uk
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ABSTRACT

**Purpose/Aims:** Patient-reported outcome measures (PROMs) can be effectively utilised to uncover the unmet needs of women with cervical cancer for supportive care. Our aim was to explore the feasibility and acceptability of PROMs-driven, nurse-led consultations to enhance delivery of supportive care to women with cervical cancer during active anti-cancer treatment.

**Design:** A two-phase, mixed-methods, prospective study was conducted. Main research variables included feasibility and acceptability parameters of the trialled intervention.

**Methods:** Pre-consultation PROM data were collected during three consecutive, monthly consultations, and used by the Gynaecology cancers nurse specialist (CNS) to deliver personalised supportive care. The Problem Checklist and Cervical Cancer Concerns Questionnaire were used to aid data collection.

**Findings:** Due to considerable recruitment challenges, a recruitment rate of 27% (3/11 patients) was achieved. Two patients completed all three study assessments. Seven in-clinic patient assessments were performed over 6 months. Study participants praised the opportunity for dedicated time for patients to raise concerns and for the CNS to provide sensitive and personalised support.

**Conclusion:** Women with cervical cancer perceive important benefits from participating in PROMs-driven, time-protected sessions with their CNS. Our findings provide tentative evidence to support the feasibility and acceptability of this intervention model, and warrant future confirmation.

**Keywords:** Patient-reported outcome measures; unmet needs; supportive care; cervical cancer; cancer nurse specialists

**Implications for practice:**

- Use of PROMs to identify the unmet needs of women with cervical cancer is acceptable, and must be implemented from the point of diagnosis.
• Cancer nurse specialists (CNS) are receptive to and able to act upon PROM information, rendering them key professionals in addressing the supportive care needs of people with cancer.

• Women with cervical cancer perceive important benefits from participating in PROM-driven, time-protected and private sessions with their CNS.
INTRODUCTION

There were close to 528,000 new cases of cervical cancer worldwide in 2012,\textsuperscript{1} but incidence rates have decreased by over 40% since the late 1970s.\textsuperscript{2} Advances in both diagnostic tests and treatments for gynaecological cancers have also led to improved survival rates, with over 60% of women now surviving to $\geq$10 years after diagnosis.\textsuperscript{2} This also means that the number of women set to deal with the aftermath consequences of cervical cancer is on the rise.\textsuperscript{3,4}

Negative effects of cervical cancer may emerge soon after diagnosis and treatment, whilst some women may continue to deal with the adverse physical, psychological and social difficulties for up to twelve months after diagnosis, and beyond.\textsuperscript{5-8} The requirement to provide on-going and comprehensive supportive care to these individuals is therefore prominent.\textsuperscript{9,10}

Developing new clinical supportive care services for women with cervical cancer means identifying ways to feasibly assess and effectively address patients’ needs. The use of patient-reported outcome measures (PROMs) may prove useful to this end. PROMs are defined as measurements of an aspect of a patient’s health status that come directly from the patient.\textsuperscript{11} The collection of PROM data in clinical practice has been reported to have a number of benefits on patient, clinical process and health service outcomes.\textsuperscript{11-13} As a result, PROMs are high on political agendas,\textsuperscript{14} as their use in informing care delivery and organisation of health services becomes apparent. Nurses’ prime position in the healthcare system means that they have a direct influence on patient experiences and outcomes of care,\textsuperscript{15} while their overall positive attitude towards collecting and utilising patient-reported health data has led to them being regarded as the most appropriate health professionals to lead on use of PROMs in clinical practice.\textsuperscript{16,17}

Currently, additional research is needed to explore feasibility/acceptability parameters around the use of PROMs by Gynaecology cancers nurse specialists (CNS) to assess and address the supportive care needs of women with cervical cancer in the acute care setting, and to evaluate how use of PROMs impacts on patient outcomes and clinical practice.\textsuperscript{18} The current study aimed to generate evidence to address this gap.
METHODS

After obtaining Research Ethics approval (13/ES/0056), we conducted a two-phase, mixed-methods exploratory study within one NHS board in Scotland.

Phase 1

Phase 1 aimed to identify what outcomes are important to women with cervical cancer and health professionals involved in their care. We therefore combined evidence from a systematic literature review with data from subsequent focus groups interviews with women with cervical cancer and health professionals involved in their care.

Systematic literature review

A systematic review of the literature aimed to appraise the empirical evidence on the supportive care needs of women with cervical cancer.19 In addition, the review aided in the identification of PROMs, developed to assess the supportive care needs of this patient population, and used in the reviewed studies. The identified PROMs were added to the pool of validated supportive care needs PROMs already known to the research group from previous reviews.20–22 They were all considered for use in Phase 2 of this project.

Stakeholder interviews

Two focus group interviews – one with patients and one with health professionals – were planned, each consisting of no more than ten participants. In focus groups, participants are guided via a facilitated discussion to express their attitudes and opinions towards a defined concept/topic, by building on each other’s ideas.23 As such, focus groups have become very useful for needs assessment and project evaluation purposes.24 Eligible patients were (a) diagnosed with cervical cancer (of any stage) within the past 12 months; (b) deemed by a member of the health team to be physically and psychologically fit to participate in the study; (c) able to read and write English; (d) able to provide written informed consent; (e) aged 18 years or over; and (f) able to provide consent for members of the research team to access their case notes. Members of the multidisciplinary team were also invited to participate in the focus group, including the Gynaecology CNS, who identified eligible patients and delivered the intervention in Phase 2.
At the end of each focus group, we involved participants in a 10-minute exercise. Copies of the previously selected PROMs were distributed to each group. We asked all participants to review the PROMs and select, in order of descending preference, the two ‘most appropriate’ for use with women with cervical cancer. Participants were asked to focus on such aspects as overall presentation, length, wording, and comprehensiveness as indicators of PROM appropriateness.

Phase 2

Phase 2 entailed a prospective, repeated-measures study that aimed to involve up to 30 women with cervical cancer as per current available guidance for early feasibility testing. Participation of the CNS was re-confirmed for Phase 2. Patient eligibility criteria were identical to those used in Phase 1. All consenting patients provided written informed consent. None of the patients who were involved in Phase 1 participated in Phase 2.

Procedures

Patients were planned to participate in Phase 2 over three, equally-spaced (monthly) time-points: baseline (within 12 months post-diagnosis) (T1); 1 month following entry into the study (T2); and 2 months following entry into the study (T3). This timeline was chosen together with focus groups participants to allow sufficient time for feasibility testing, whilst minimising the attrition rate.

All eligible patients were thoroughly informed about the purposes and procedures of the study, and provided written informed consent. At each time-point, participating patients were booked on an appointment with their Gynaecology CNS. Whilst in the clinic and prior to their consultation with the CNS, patients were asked to complete the set of selected PROMs selected in a quiet room. Subsequently, the Gynaecology CNS met with the patient and used the information collected via the PROMs to direct consultations, identify the patient’s supportive care needs, and intervene accordingly. The CNS documented the supportive care needs she identified during the consultation and any resulting interventions in author-developed case-report forms. Up to ten participating patients and the Gynaecology CNS were planned to participate in one-to-one, end-of-study interviews to explore their perceptions on the use of PROMs in clinical practice.
Data analysis

In light of poor recruitment in Phase 2 (see Results), the initial analysis plan was revised and a case-report analysis approach was pursued. Pseudonyms were used for presentation purposes. PROM data were entered in Microsoft Excel spreadsheets and analysed using descriptive statistics and graphs/radar plots for each case report. Frequency counts for each response were generated to describe response patterns for PROM domain and domain scores, and quantify missing data. Focus groups and end-of-study interviews were audio-recorded and transcribed verbatim. Thematic analysis was used to identify, analyse and report patterns within interview data.26

RESULTS

Phase 1

Structured literature review

Dealing with fear of cancer recurrence, concerns about appearance/body image, lack of sexual desire, requiring more sexuality-related information, dealing with pain, and dealing with difficulties in relationship with partner were the most frequently cited individual needs (≥4 studies).19 Based on this evidence and drawing on our database of supportive care needs PROMs, we concluded that the following eight supportive care needs PROMs would be discussed in subsequent focus groups:

Supportive Care Needs Survey – Short Form 34 (SCNS-SF34);27 Problems Checklist;28 Cancer Needs Questionnaire – Short Form;29 Psychosocial Needs Inventory;30 Comprehensive Needs Assessment Tool in Cancer (CNAT);31 Cervical Cancer Concerns Questionnaire (CCCQ);32 Cancer Rehabilitation Evaluation System – Short Form;33 and Cancer Needs Distress Inventory.34 These PROMs were selected because they are relevant, brief and, in combination, cover the needs of women with cervical cancer. Only one of them was specifically developed/adapted for use with this patient population.

Stakeholder interviews

The first focus group involved four women with cervical cancer. Two women were in their 30s and two in their 50s (age range 35-55). Three women were in full-time employment, whereas one was unemployed. None of the women was married or partnered. Initially, the discussion focussed on the diagnosis of cervical cancer and how different people react to the diagnosis. One woman indicated that she took in all the cancer information she could, whereas another one did not want as much
information. All women felt it was the manner in which health professionals conveyed information to
them (i.e. “very matter of fact”, “not sugar-coated”) that was of particular importance.

The group also discussed their experiences of chemotherapy, revealing a range of symptoms
that included constipation, fatigue, pins and needles, sleep disruption and flatulence. Notably, all
women felt that during treatment it was good to be among people, who were experiencing the same
thing as they were. This was in contrast to their family and friends “who do not really know what it is
like”. These women did not feel the need for spiritual or pastoral care, but they admitted to not being
particularly religious. The suggestion of discussing spiritual needs was initially interpreted by one
patient as dealing with death, for which there was an obvious fear. Finally, when discussing if their
needs had been covered during consultations, one woman noted that she was never asked if she
wanted to preserve her ovaries, and stressed the importance of fertility concerns being identified and
addressed.

The second focus group involved three health professionals (1 Gynaecology CNS and 2
consultant oncologists). Two health professionals had over 10 years of experience working with
women with cervical cancer. The main message from the health professional focus group was the
need to have individualised assessments, given the mix of patients health professionals normally deal
with. Various demographic and clinical characteristics (e.g. patients’ age, where they are in their
reproductive cycle, stage of cancer, and type of treatment) have to be taken into account.

The health professionals indicated that, although they would normally suggest to women with
cervical cancer to make use of available community services to get additional information and
support, one-to-one meetings were felt as more protective of patients’ privacy and conducive to open
discussions about the effects of cervical cancer. The group felt that no set of needs should be given
priority over any other for, if it is truly patient-centred care, then it should be guided by each patient’s
own priorities.

Finally, when asked if they would like a formal way to assess to supportive care needs, one
consultant was somewhat sceptical. In contrast, the other two health professionals seemed more open
to introducing formal needs assessments in clinical practice. The CNS reported that she had used the
Distress Thermometer in the past and found it “very valuable”. The second consultant highlighted
how important it would be to have a CNS in the consultation room every time there is a new or a
recurring diagnosis to ensure that women’s needs are identified and addressed.

**Decisions made prior to Phase 2**

The health professionals chose the Problem Checklist as the most acceptable PROM in terms of
presentation and wording, whereas patient participants selected the CCCQ. Taking both perspectives
into consideration as well as the need for comprehensive assessments, a final decision was made to
use both the CCCQ and the Problem Checklist in Phase 2. One item was added to the Problem
Checklist, pertaining women’s concerns about their ability to have children/become pregnant in the
future, in line with findings from Phase 1.

The CCCQ is a 37-item questionnaire that aims to assess women’s concerns in relation to
cervical cancer and its treatment. A combination of Likert-type or numerical scales, and open-ended
questions are employed. Thirty-two items are used to categorised into seven domains of concern,
including (1) communication with the treatment team, (2) treatment issues, (3) sexuality, (4)
prognosis, (5) cause/transmission risk, (6) partner relationship, and (7) relationship with others. Each
item is rated for level of concern during the past week on a seven-point Likert scale, ranging from 1
“not very much” to 7 “very much”.

The Problem Checklist assesses the extent of patients’ concerns or difficulties in each of 16
aspects of their lives as a result of cancer and/or its treatment. Items are scored on a four-point scale
(1 “no difficulty” – 3 “severe difficulty”). In our study, a 17-item checklist was used.

**Phase 2**

**Feasibility and acceptability estimates**

Between July 2014 and July 2015, recruitment in Phase 2 had to be postponed twice due to
considerable challenges with patient availability for a focus group, and a period of clinician absence
from work. In the limited amount of time left (July to October 2015), 11 eligible women with cervical
cancer were invited to the study. Four women declined participation, due to lack of time, deteriorating
physical condition, or lack of interest. In addition, four women never contacted the CNS or attended
the clinic after being invited, thus lack of interest was also assumed. The final sample consisted of 3
women, who provided written informed consent.
A recruitment rate of 27% (3/11) and an average recruitment pace of 1 participant per month were achieved. Two patients (67%) completed all 3 study assessments, with one patient withdrawing soon after baseline assessment due to declining health status.

The Gynaecology CNS performed a total of 7 in-clinic patient assessments within a period of 6 months (i.e. the period when the study was ‘open’ for recruitment and follow-up). Full documentation records were received for each in-clinic assessment. Reflection questions were filled out for all 7 in-clinic assessments. Completeness of background data was 95%.

In terms of data collection, 7 questionnaire packs were returned (100%), one for each-clinic assessment. Data completeness analysis indicated that across 378 actual questionnaire data, only 0.8% were missing across 3 assessment points. Questionnaire completeness reached 99.5% at baseline, and remained high at T2 and T3 (99.6% and 99.8%, respectively).

**Prevalence and over-time change in patients’ needs**

**Case #1: Kristie** – Kristie was a single woman in her 40s when she was diagnosed with stage 2b cervical cancer. She was employed, but on sick leave. She reported no comorbid illnesses. At the time of her baseline assessment, she was receiving a combination of chemo-radiation treatment with brachytherapy. Her performance status was assessed as moderate, meaning that she was ambulatory and capable of all self-care, but unable to carry out any work activities.

At baseline, Kristie’s concerns revolved around receiving clear information about her diagnosis and treatment, coping with chemotherapy and related fatigue, surviving cervical cancer, and dealing with the possibility of a cancer recurrence and the effects of cancer on her family and friends. The latter issues seemed to abate during the second assessment (T2), but the same treatment-related concerns were still prominent as she went through full cycles of active anticancer treatment, and her physical functioning deteriorated. At T3, when Kristie was close to finishing with treatment, her concerns were considerably lower and she appeared to be in control of even the most consistent ones, i.e. surviving cancer and fearing of cancer recurrence. The radar plot of CCCQ domains of concern supports this trend (Figure 1a).

Answers on the Problem Checklist also revealed moderate to severe difficulty with dealing with work-related and financial matters throughout the study, but mainly at T2 (Figure 1b). During her
appointments with the CNS, Kristie specifically requested advice about symptom management, information about brachytherapy, and help to deal with feelings of isolation, fears of cancer recurrence, financial concerns, an altered body image, relationships, menopause, and current and future work issues.

**Case #2: Anna** – Anna was a young, married woman and a mother of two. She was a full-time employee, currently on sick leave. She was diagnosed with stage 1b cervical cancer and planned to receive post-surgery chemo-radiation therapy combined with brachytherapy. She reported no comorbid medical conditions. Her baseline performance status was assessed as moderate.

Regardless of time-point of assessment, Anna appeared to be overly distressed and in need for on-going help and support. Her responses on the CCCQ (scoring ‘7’ for 99% of all items and assessments) were indicative of a person overwhelmed by the diagnosis of cancer – especially at this very young age – that struggled to come to terms with the illness and its treatment, and the effects of cancer on her, her family and their future (Figure 2a).

A similar picture was obtained from reviewing Anna’s responses on the Problem Checklist. Practical, daily living and social needs were evident especially as Anna moved on with her treatment (Figure 2b). The only area that was of no concern for her was her ability to have children in the future. During consultations, Anna was able to ask for help to deal with the “information overload” about her diagnosis and treatment. In addition, she felt the need to discuss issues around the effects of cancer on her relationship with her husband, her fears about dying and about the cancer returning, her feelings of being abandoned by friends and family, financial difficulties, and her efforts to accept that she could not be her children’s caregiver anymore and had to rely on her own parents.

**Case #3: Ruth** – Our third participant, Ruth, was a 30-year-old married woman, on extended sick leave after her diagnosis. Ruth reported no comorbid illnesses, while her baseline functional status was good. Of note, Ruth was diagnosed at an advanced stage and the aggressiveness of her tumour increased her odds for a poorer clinical outcome. Ruth was put on a combination of chemo-radiation therapy with brachytherapy, but eventually, her rapidly deteriorating condition prevented her from completing the study. She dropped out soon after her first consultation.
Although a comparative, over-time analysis of Ruth’s data was not possible, her baseline assessment revealed a multitude of concerns and needs (Figure 3a). This was particularly in relation to the short- and long-term effects of treatment, survival, sexuality and intimacy issues, the impact of cancer on her relationship with her husband and her family and friends, and even the possibility of her being stigmatised for having a cervical cancer diagnosis. Ruth’s Problem Checklist also confirmed her difficulty to come to terms with her illness and deal with such issues as relationships and sexuality.

Additional areas of concern included work, finances, managing at home, and keeping up with her interests (Figure 3b). However, Ruth’s primary difficulty was that of coming to terms with the possibility of her not being able to have children in the future. In her questionnaire, she scored this as a severe difficulty, underlined the relevant item twice, and wrote: “Side effects on young women with no children. Losing the right to become parents (support)!!” Her consultation appointment focussed on these areas.

**End-of-study interviews: Kristie**

In her exit interview, Kristie focussed on how the PROM intervention helped her flag needs or concerns that were initially less obvious to her: “Well when I was filling it [the questionnaire] in, it was things that I never really thought of... you think, “I probably did need more information.” During the PROM-driven consultation, Kristie got clearer and more personalised information on a number of pertinent issues, predominantly those practical, daily living and family-related ones. The secured time with the CNS was perceived as beneficial; a useful adjunct to the support she was getting from her family: “…cause it’s somebody [the CNS] there to sit and listen to your problems. Somebody that was out [with] the family.” During her consultations, Kristie opted for a family member to attend, too, possibly as a way to feel more comfortable and secure.

Overall, the PROMs were found to be clear enough and straightforward. Kristie stressed that the wording of questions that patients are asked to respond to need to be as clear as possible to prevent confusion: “Don’t put like big words in, just put simple questions just to ask them.” Kristie also mentioned that she would welcome more time with the CNS if that was feasible. Getting access to such a service from the point of diagnosis onwards was also flagged as very important: “At the beginning. Mhmm. Definitely… To let people know they’re not alone... But I think maybe once you
“get the first, maybe 6 months maybe a year out you, I think I’d probably feel better. I wouldn’t be worrying it as much.” Kristie felt that, after treatment, she was struggling to communicate her needs to health professionals, especially those seen at the community (GP or district nurse), who had limited insight in her condition and experiences. Relaying PROM information to GPs or district nurses was seen as an effective way to bridge the gap: “So they’ve got a, more understanding of what folk are going through. Cause every time I’ve went to them, it’s like, ‘And what is it that’s wrong with you?’”

End-of-study interviews: Anna

Anna spoke about how completing the PROM and discussing flagged needs with the CNS was helpful as she was trying to cope with her diagnosis: “It’s just obviously when I came to complete [the questionnaire] with [the CNS] I could kinda speak to her. And speak over all the, the questions and she would kinda help me kinda answer it.” Using the PROM gave Anna the opportunity to disclose sensitive information and subsequently discuss it with the CNS in a non-judgmental environment, whilst it prompted her to seek help for any issues rather than ignore them: “I think [it was] just because of the questionnaire [that helped to bring up the issue]. ...I had some bleeding after sex the other day. ...I would [...] just have shrugged that off. But obviously because of what I’ve been through now, I think ‘I need to ask somebody’.”

Anna stressed that the one-to-one and face-to-face approach was an important component of the intervention. She also pointed out that appropriate timing is important, and that the intervention should be placed at the very start of the journey and then during staging, because: “…it’s the times in-between that you need somebody to explain.” Anna suggested that allowing one to write a short paragraph about themselves would help put the needs assessment into perspective and further tailor the advice to the specific person: “I don’t know if you could write a wee bit about yourself. And what, that’s happened to you. Kinda would maybe be a bit helpful.” Moreover, involving another person with a similar experience was seen as a way to further improve the quality of the support given: “I think it would be beneficial to people to kinda maybe speak to somebody who’s been through, been through all the treatment. Having somebody there that can relate to what you’re going through”

Finally, Anna agreed that making PROM data available to GPs would be a helpful strategy to enhance receipt of personalised support in the community.
End-of-study interviews: Gynaecology CNS

The CNS felt that the PROMs helped patients to feel more comfortable and open up, which facilitated assessment of a wide range of needs. Using the PROMs gave the consultation structure and improved the quality of the discussion: “I do discuss everything with these women anyway, but I felt that, that the questionnaire, it was a bit more structured and it was, like I used it as a guide to me, to keep me in the track...” Eventually, the structure that the PROMs instilled facilitated a patient-led consultation, which “was very useful... [and it] gave them the choice... and it was their choice what they wanted to discuss and what was a priority at that particular time.” The CNS felt that the PROMs were overall “very comprehensive and there was absolutely nothing at all missing...”, but there were certain areas that appeared more challenging for patients to raise (“intimacy...they [the patients] were a wee bit apprehensive about discussing it”) or for the CNS to handle (“there was one [area] that I found quite challenging; death and dying”). Did the CNS have adequate resources to respond to such patients’ needs? “I think you know, you, you have to work with the resources that you’ve got and if I felt that I couldn’t have dealt wi’ these women then obviously I would have signposted them to other agencies.” Eventually, the CNS seemed clearly satisfied with being involved in the study: “I thoroughly enjoyed doing that study... because I think, you know, that the information that we get from this could generate a change”.

DISCUSSION

Our systematic review identified 13 studies, demonstrating the variability and extent of unmet needs of women with cervical cancer across different phases of the illness trajectory. Yet, interventional research to develop and evaluate strategies to address these needs is lacking. This small study has shown that nurse-led, PROMs-driven consultations to identify and address the supportive care needs of women with cervical cancer are acceptable to and considered worthwhile by both care recipients and care providers. Nonetheless, certain feasibility parameters need to be taken into careful consideration before the intervention is deployed in clinical practice. For instance, given the unsatisfactory recruitment rates, it remains unclear whether the intervention itself was of no interest to possible candidates or whether it was affected by how the intervention was delivered (needs assessment sessions on top of normal in-clinic consultations) or how women were approached to
participate (letters sent to eligible women and a mixed opt-in/opt-out method was applied). With only one new attendee per month, the time-effectiveness of the intervention is also unclear. Adopting a more flexible schedule for delivery of the intervention, whereby PROMs-driven consultations coincide with pre-arranged hospital visits and/or are delivered via telephone or online, could further encourage participation.

Unlike researcher-supported studies, here we relied on an actual member of the clinical team to incorporate the intervention and research activities in her workload. This approach renders our findings significant and relevant to clinical practice as we were able to establish a realistic view of the facilitators and barriers of implementing this intervention. That said, it was made apparent that single-handedly delivering a time-intensive intervention may hinder adequate testing and implementation. It is thus important to identify ways to either bring the intervention down to delivery schedule that is more manageable for lone providers or ensure adequate nursing support.

The unique nature of the consultations (one-to-one, face-to-face, patient-driven, and time-protected) was highly praised by both intervention recipients and provider as it fostered a secure place for women with cervical cancer to disclose intimate and/or hard-to-verbalise issues. Both Kristie and Anna endorsed the standardised use of an easy-to-understand supportive care need PROM as a means to help them shortlist, report and prioritise their needs. Although we did combine two PROMs, totalling 54 items, we received no complaints regarding time or length of the assessment. It is worth noting that in Phase 1, health professionals opted for brevity in needs assessments, whereas patients focussed more on relevance and comprehensiveness. In Phase 2, we decided to prioritise patients’ preferences, but we do appreciate the need to find ways to minimise clinical work overload. If this model is to be implemented in practice, it will be an interesting future step to explore whether use of a bespoke and concise, yet equally comprehensive, needs assessment PROM could be used, and/or additional clinical resources become available.

In exit interviews, both women agreed that timing of the intervention was appropriate and relevant, but specifically flagged the post-diagnosis and post-treatment period as the ones where greatest support is necessary. Interestingly, both women also mentioned how helpful it would be if summaries of their needs were also shared with GPs and community nurses in order to increase
understanding of their situation and facilitate more tailored discussions in the community. The need to
attend to cancer survivors’ rehabilitation needs is a known one, and major policy documents
advocate development of mechanisms to promote clinical continuity and better manage transitions of
care. Previous research has shown that while physical concerns are often addressed by the GP
and/or community nursing staff, psychosocial aspects (such as fear of relapse or social adjustment) are
often under-reported and under-assessed, and thus neglected. Our findings pose a clear indication for
needs assessment data to be shared with all health professionals involved in the provision of care to
(at least) women with cervical cancer in order to enable effective communication that can lead to
seamless care.

From a clinical point of view, the Gynaecology CNS perceived engagement in the collection
and use of patient-reported data as an enlightening and educative activity. In Phase 1, it was
interesting to see how one consultant argued against the ‘mechanistic’ nature of PROMs-driven
assessments, essentially opting for needs assessments that are based on individual clinical expertise
and experience only. Conversely, existing evidence is largely supportive of a combination of
structured (PROMs-based) assessments and patient management that is based on clinical expertise
and specialised training to ensure that the holistic care that patients expect to receive is indeed
provided. PROMs can be the means to unveil unmet needs that can clinical expertise can help to
address. It can’t be ignored however the possibility for some concerns to be difficult to explore or
handle. In our study, concerns about death and dying proved challenging to address, whilst
assessment of intimacy issues was met with apprehension. As part of a needs assessment intervention,
nurse specialists could be trained to provide education for possible patient adjustment issues or
address women’s sexuality needs, whilst clear routes of referral could enable timely and appropriate
referral to clinical specialists.

As yet, we cannot reliably estimate the potential effectiveness of the intervention nor can we
claim superiority of consecutive, monthly assessments over one-off, post-diagnosis assessments or
assessments timed in line with major patient transitions. Provided that such aspects are clarified in a
future replication study, this intervention could be an effective means for Gynaecology CNS to
provide comprehensive, nurse-led supportive care to women with cervical cancer.
STRENGTHS AND LIMITATIONS

We thoroughly reviewed the existent literature and subsequently engaged patients and health professionals in the actual planning of the study. This phased approach helped us to customise and refine aspects of the intervention in an attempt to meet users’ preferences, expectations and priorities, and increase the intervention’s feasibility and acceptability. Subsequently, we relied on a set of well-validated PROMs to collect information in a reliable and comprehensive way. We then employed different sources of information to comprehensively investigate the study’s feasibility and acceptability, including observation, questionnaire and interview data. Last, evaluation of the intervention with minimal research support and clinical practice assimilation conditions allowed for a realistic assessment.

The study should nonetheless be interpreted in the context of a number of key limitations. Neither PROM completion nor consultation appointments were timed; however, none of the participants reported the intervention as time-consuming. With a small sample size like this, whether reliable feasibility estimates were obtained is unclear as is the influence of demographic/clinical characteristics as moderators of feasibility. Relatedly, we were unable to provide estimates of responsiveness to change or effect sizes for any of the intervention PROMs. Finally, this was a single-centre study, thus reflecting current facilitators and barriers in the implementation of PROMs-driven supportive care intervention for women with cervical cancer within the participating NHS board. Whether feasibility and acceptability of the intervention is similar in diverse clinical contexts remains unknown.

CONCLUSIONS

Testing the use of PROMs by nurse specialists in the delivery of supportive care to women with cervical cancer indicated that this approach appears to be acceptable, but its feasibility requires further evaluation. Congruent with the literature, we confirmed that CNS are key professionals in the delivery of supportive care, and are receptive to and able to act upon information gleaned from supportive care needs PROMs in clinical practice. Women with cervical cancer perceive important benefits from participating in PROMs-driven, time-protected and private sessions with their CNS. Nevertheless, our
findings provide only tentative evidence to support the future use of PROMs as part of nurse-led consultations in this area, and warrant further confirmation in the future.
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Conflicts of interest

The authors declare that there are no conflicts of interest, financial or otherwise.

Statement of Authorship

All authors have equally contributed to the preparation of this manuscript.
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Figure Legends

**Figure 1.** (A) Kristie’s CCCQ radar plot showing scores on seven domains of concern over three time-points. Higher scores indicate greater concerns. (B) Kristie’s responses on the Problem Checklist over three time-points.

**Figure 2.** (A) Anna’s CCCQ radar plot showing scores on seven domains of concern over three time-points. Higher scores indicate greater concerns. (B) Anna’s responses on the Problem Checklist over three-time points.

**Figure 3.** (A) Ruth’s CCCQ radar plot showing scores on seven domains of concern for T1. Higher scores indicate greater concerns. (B) Ruth’s responses on the Problem Checklist at T1.
Figure 2

(A) Radar chart showing the communication with the clinical team, treatment, sexuality, relationship with partner, causes/transmission, and prognosis for T1, T2, and T3.

(B) Bar chart showing the difficulty with various aspects such as coming to terms with illness, coping with treatment, getting on with hospital staff, work, finances, managing at home (chores), managing at home (self-care), getting on with family, relationship with partner, sexual relationship, coping with children, social life, interests, mood, religious beliefs, thinking/memory, and ability to have children for T1, T2, and T3.