Improving quality of life in cancer patients through higher participation and health literacy: study protocol for evaluating the oncological social care project (OSCAR)

Johann Frick 1*, Daniel Schindel 1, Pimrapat Gebert 1,2,3, Ulrike Grittner 2,3 and Liane Schenk 1

Abstract

Background: Cancer patients experience psychological and social distress due to their medical treatment and social issues. However, continuous and specialized social support is still lacking. In Germany, a group of company health insurance funds has developed an approach to support cancer patients with monthly structured interviews conducted by specially trained Social Care Nurses. The nurses will identify patient needs in order to provide help with medical, personal, and social matters. One aim of the scientific evaluation is to analyze the effect of the consultations on various patient-reported outcomes, especially quality of life. The evaluation concept will be described in this study protocol.

Methods/design: The evaluation is a non-randomized, controlled, multi-center intervention study with a mixed-method design. It consists of three research modules which include primary data from questionnaires, and claims data from the health insurance funds. In Module 1, cancer patients will be recruited to form an intervention group (OSCAR, \( n = 150 \)) and a control group (\( n = 200 \)) in four study centers for a period of 1 year. One baseline and three follow-up questionnaires will be conducted to survey the patient-reported outcomes. Relevant secondary outcomes are health literacy, participation, and physician-patient communication. In Module 2, claims data will be used to analyze cost effects and thereby assess effectivity and hospitalization. Module 3 will involve a qualitative analysis of project diaries kept by the Social Care Nurses. The diaries will record the nurses’ practical experiences and the benefits of deploying OSCAR across the German healthcare system.

Discussion: OSCAR is an innovative way of providing cancer patients with continuous support to improve their quality of life. The evaluation concept aims to assess the effects of the monthly consultations by the Social Care Nurses on the patients, and will use a mixed-method design. The results are important for assessing the transferability of OSCAR to the healthcare system as a whole.

Trial registration: German Clinical Trials Register (DRKS-ID: DRKS00013640). Registered 29 December 2017.

Keywords: Social support, Cancer, Quality of life, Patient preferences, Health literacy

* Correspondence: johann.frick@charite.de

1Institute of Medical Sociology and Rehabilitation Science, Charité – Universitätsmedizin Berlin, Campus Charité Mitte, Charitéplatz 1, 10117 Berlin, Germany

© The Author(s). 2019 Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The Creative Commons Public Domain Dedication waiver (http://creativecommons.org/publicdomain/zero/1.0/) applies to the data made available in this article, unless otherwise stated.
Background

Demand for social support for cancer patients

Cancer causes a high number of incident cases in Germany every year (229,920 women and 252,550 men in 2013) [1]. Due to the progression of the diseases, patients and their relatives experience distress caused by medical treatment and social issues [2, 3]. The misunderstanding of the disease can lead to inappropriate self-assessments of one's prognosis and a choice of unsuitable medical treatments [4]. Simultaneously, patients receive overtreatment even though they have an incurable disease [5]. In these situations, palliative care could be appropriate for improving quality of life [6]. Enhancing palliative care can help patients to find suitable medical treatments that can increase quality of life. However, the specific needs of cancer patients are not taken into account by the healthcare system because of a lack of palliative care [7]. Currently, patient navigation programs are used to help patients with medical, nursing, psychological, and healthcare-related issues. Recent studies have attempted to evaluate the impact of these programs (such as Onkolotse by the Saxon Cancer Society) [8, 9]. In this context, patient-reported outcomes (PROs) are very important for examining the subjective benefit of interventions from the patient's perspective, with the focus on relevant outcomes (e.g. quality of life) [10].

Development of the OSCAR project

The Oncological Social Care Project (OSCAR) was developed by the German company health insurance fund Pronova BKK. The concept is based on the Saxon Cancer Society’s Onkolotse navigation program for cancer patients. An essential component of OSCAR is the additional social support provided by the Social Care Nurses (SCNs), who aim to identify deficits in healthcare utilization by cancer patients and support them in relevant areas (e.g. the medical treatment of the cancer, psychosocial support for anxiety, practical tips regarding social security services, and organization of rehabilitation). Moreover, the SCNs are regular and accessible contact partners in a complex and fragmented multidisciplinary treatment process. Each patient is accompanied for 1 year, regardless of whether they are being treated as inpatients or outpatients, or are currently not receiving any therapy. The intervention aims to access and improve the quality of life for patients with advanced cancer and a poor prognosis. It uses twelve structured interviews over the course of 1 year. The SCNs received training from the Saxon Cancer Society. OSCAR is implemented by five SCNs at four locations.

Methods

The OSCAR evaluation is a mixed-method study consisting of three research modules. Modules 1 and 2 involve a non-randomized, controlled, multi-center intervention study which compares primary data from regularly conducted questionnaires and claims data from German statutory health insurance funds in the intervention and control groups. In addition, a qualitative content analysis is planned in module 3 and will be based on the project diaries completed by each SCN. The analysis will assess OSCAR’s effectiveness and the benefit of implementing it in the German healthcare system (Fig. 1).

Study objectives

Primary objective:

To compare quality of life over study period between the intervention group (patients who had met an SCN) and the control group (Module 1).

Secondary objective:

1. To compare the health literacy, participation, and physician-patient communication in the intervention and control groups (Module 1)
2. To explore the associations between quality of life and health literacy, participation, and physician-patient communication (Module 1)
3. To compare the incidence rate of outpatient visits in both groups (Modules 1 and 2)
4. To compare the healthcare costs in both groups (Module 2)
5. To assess the SCN’s practical experiences (Module 3)

Inclusion and exclusion criteria for participation in the evaluation survey in modules 1 and 2

Inclusion criteria:

- Aged ≥18 years
- At least a combination of one type of cancer and one operation and procedure, as defined by the ICD and OPS codes listed in Additional file 1

Exclusion criteria:

- Advanced dementia
- Acute addictions

Module 1: primary data, patient-reported outcomes

Participant recruitment

The recruitment was carried out in four different hospitals with oncology departments across three cities in two states in Germany. Two of the hospitals are located in Berlin, while the other two are in Leverkusen and Duisburg.

In all four study sites, the SCNs recruited BKK-insured patients for the intervention group by obtaining informed consent. The SCNs are responsible for the monthly consultations to support the patients in the
intervention group. Patients who are insured with a different provider were recruited by Study Nurses (SNs) for the control group. The SNs’ main task is to conduct the evaluation questionnaires in both study groups up to four times within 1 year. The evaluation’s baseline survey will be conducted after recruitment (t0) and followed up after 90 (t1), 180 (t2), and 365 (t3) days. The patients can choose between personal, telephone, and postal interviews, depending on their health status and which method is most comfortable for them. Dividing responsibilities between SCNs and SNs is necessary to ensure the independence of the evaluation surveys. Patients in both groups were recruited at the same study sites between February 2018 and the end of February 2019.

**Questionnaire**

The items for the evaluation questionnaire are listed in Table 1:

Further instruments are used to measure sociodemographic variables, especially with regard to age, sex, migration status, and social status [19–21]. Finally, participants in the intervention group will be asked to evaluate their...
Table 1 Items in the evaluation questionnaire

| Item                                                                 | Description                                                                 |
|----------------------------------------------------------------------|-----------------------------------------------------------------------------|
| Quality of Life of Cancer Patients (EORTC QLQ-C30) [11]              | The EORTC QLQ-C30 is a questionnaire for measuring health-related quality of life, and is provided by the European Organisation for Research and Treatment of Cancer. Patients are asked to answer 30 questions, which address e.g. physical, emotional, cognitive, and social issues. 28 items have four response options. Two items for the global health status and the quality of life have a seven-point scale. The EORTC QLQ-C30 consists of five functional scales and three symptom scales. In addition, a global health status (QoL) scale as well as six single items. All of the scales and the single items will use a linear transformation to standardize the raw score. The scores range from 0 to 100. |
| Illness Perception Questionnaire (IPQ) [12]                         | The IPQ consists of 64 items on 8 scales, and measures the individual’s beliefs and feeling about their illness. The illness coherence subscale consists of five items and is used for the evaluation questionnaire in this study. The scores will be summed and divided by the number of items. |
| Patient Reaction Assessment (PRA-D) [13]                            | The PRA-D is a self-description instrument to record the patient-perceived quality of the relationship between patients and physicians. In this study, we adapted five communication questions to a five-point Likert scale (from 1 = "I strongly disagree" to 5 = "I strongly agree"). |
| German modified version of the Autonomy Preference Index (API-Dm) [14]| The API-Dm measures patients’ preferences for participation and information regarding medical decisions. Patients are asked to answer eleven questions using a four-point Likert scale (from 0 = "strongly disagree" to 4 = "strongly agree"). The information preference consists of seven items, with scores ranging from 0 to 28. The decision preference consists of four items, with scores ranging from 0 to 16. The total score of each patients’ preferences will be transformed into scores ranging from 0 to 100. |
| Decisional Conflict Scale (DCS-10) [15]                            | The DCS-10 is a self-reported questionnaire, which allows an evaluation of decision conflicts among patients. The short version of the DCS comprises ten items grouped into four subscales: uncertainty, information, values clarity, and support decision. Each item is measured on a three-point Likert scale (0 = “yes”, 2 = “unsure”, 4 = “no”). Scores range from 0 (no decision conflict) to 100 (extremely high decision conflict). |
| European Health Literacy Survey (HLS-EU-Q6) [16]                    | The HLS-EU-Q6 is the short version of the 47-question European Health Literacy Survey. It is divided into three areas: healthcare, disease prevention, and health promotion. Each of the six questions has five possible responses. A mean score is calculated and allocated to one of three categories: insufficient health literacy (range 1–2), problematic health literacy (2–3), and sufficient health literacy (3–4). |
| Medical Care Utilization [17]                                       | Medical care will be assessed by the frequency of utilization of different therapies and physician groups. |
| Oslo Social Support Scale (OSSS-3) [18]                             | The OSSS-3 is a brief measurement of social support. It consists of three questions that address the number of close people, interest and concern from other people, and practical help from neighbors. The scores range from 3 to 14 and will be categorized into three groups: 3–8 indicates poor support, 9–11 shows moderate support, and 12–14 reflects strong support. |

Contact with the SCN for the last 3 months. The items are related to specific consultation topics and whether or not the support was useful for the respective topic. Participants will also be asked about the quality and quantity of contact with the SCN.

Statistics

Sample size calculation

A sample size of 100 participants in the intervention group and 150 in the control group will reach a power of 80% at a two-sided level of significance of 5% to detect a difference in the EORTC QLQ-C30 (scores range from 0 to 100) with a Cohen's d effect size of 0.4 (mean difference of ten scores and standard deviation (SD) of 25 scores) [11]. Given that the effect size might be smaller due to a lower mean difference or higher SD, and that the severity of cancer can cause higher dropout rates, we aim to include 150 participants in the intervention group and 200 in the control group (so a total sample size of 350 participants).

The sample size calculation based on the t-test was used in spite of the intended analysis with a baseline-adjusted repeated-measures linear mixed model (ANCOVA, three-level random intercept model to account for repeated measures in patients and clustering in centers). It can be shown [22] that a conservative approach for estimating sample sizes has the same power as a t-test with n subjects where p is the variance deflation factor, calculated by the correlation of baseline and follow-up measures. Assuming the worst case of p = 0 leads to the sample size based on the t-test.

Statistical analysis plan

All statistical tests are performed using Stata IC15 (StataCorp, 2017, College Station, TX, USA). The primary hypothesis will be tested at a two-sided significance level
of $\alpha = 0.05$. All secondary hypotheses will be tested within an exploratory framework.

Descriptive statistics and the number of participants reflected in the calculation ($n$) will be presented in each group. For continuous variables, mean with SD for normal distribution and median with interquartile range (IQR) for other distribution variables will be presented. For categorical data, frequencies and percentages will be displayed for each category. Graphic methods such as box plots and line graphs will be used for visualizing the data.

The comparison of EORTC QLQ-C30 (global health status as a primary outcome), physician-patient communication (scores), health competence (scores), participation (scores), and knowledge (scores) over the study period between the groups will be reported as mean and 95% CI, and performed using a linear mixed model (LMM) with three levels over all available time points. Random intercepts for the patient ID and for the study clinics are included in the model to account for the cluster structure of the data. To keep selection bias to a minimum, we will develop a propensity score for adjusting baseline factors. The propensity score will be used with the inverse probability of treatment weighting (IPTW) method because the results from this method are similar to a randomized trial [23, 24].

The number of outpatient visits will be counted and the study time will be calculated over the follow-up time for each patient. The incidence rate of outpatient visits per person-time will be presented, and Poisson regression will be used to compare the incidence rate between groups.

All outcomes will be analyzed using modified intent-to-treat populations including all subjects who receive at least one consultation from an SCN and for whom at least the first follow-up assessment at Visit 1 (at 3 months) is available.

**Dropouts and missing data**

Reasons for dropouts will be documented and reported. If patients are alive and missings are assumed to be missing at random (MAR), we will use multiple imputation methods based on ten imputed data sets [25] with multiple imputation by chained equations (MICE).

**Module 2: secondary data, claims data from health insurance funds**

**Design and participants**

Another part of the OSCAR evaluation involves analyzing claims data from the BKK health insurance funds. In contrast to the patient-reported parameters in Module 1, the claims data will make it possible to analyze objective parameters. This kind of data is a valid and relevant source, and has been used in many healthcare research projects in Germany. The data from Modules 1 and 2 will not be linked. If available, the secondary data will be analyzed for the period up to 12 months before and after enrollment.

**Inclusion and exclusion criteria**

This data is only available for the patients in the intervention group, who have health insurance with BKK ($n = 150$). A new, independent and anonymous control group will therefore be drawn from BKK-insured patients who did not take part in the OSCAR intervention ($n = 600$). All included patients must meet the inclusion and exclusion criteria mentioned above. Furthermore, the patients in the control group should be treated in hospitals comparable to those where the patients in the intervention group were treated. In addition, the control group will be matched with the intervention group by age, gender, and diagnosis.

**Variables**

The following aspects are of interest: time-to-death, all costs of treatment, and the incidence rate of hospitalization and outpatient visits.

**Statistical analysis plan**

Paired t-test or Wilcoxon signed rank test will be performed to compare the costs of treatment per month. The generalized estimating equation (GEE) for Poisson regression will be applied to compare the incidence rate of hospitalization and outpatient visits between intervention and matched controls. Kaplan-Meier curve and Cox regression analysis will be performed to compare time-to-death between the groups.

**Module 3: qualitative analyses of SCN project diaries**

**Design and participants**

The qualitative analyses of project diaries provide important insights into the work of the SCNs. The nurses will be asked to record their feedback concerning positive aspects of the OSCAR program, as well as opportunities for further improvements with regard to the transferability of OSCAR to the regular healthcare system. For each patient, the SCNs will record the number of additional consultations between two planned monthly consultations, and any recommendations to visit partners in their supply network.

**Analysis plan**

The following aspects are of interest for the qualitative content analysis: acceptance of OSCAR among the patients and among the participating hospitals; number of additional consultations for patients; subjective feedback and reasons for early program termination by the patient; barriers to consultations.
Discussion
This study aims to evaluate whether the OSCAR program can improve quality of life for patients with advanced cancers and a poor prognosis by enhancing health literacy and participation in therapy planning through continuous supportive care from an SCN. A variety of approaches currently exist for developing and evaluating navigation programs for cancer patients. One example is the Onkolotse project from the Saxon Cancer Society. It involves certified nurses, psychologists, and social workers, who having contact with cancer patients about 20 times within 1 year. The evaluation of this project focuses on the number of hospital admissions and psychological stress [8]. Nurses from the non-profit association Group Health contact patients weekly via telephone, and meet with them face-to-face at least once [26]. The nurses aim to develop strategies that address distress and focus on quality of life. In another project, experienced nurses attempt to identify patient needs with three different assessment tools and personal conversations [27]. The endpoints in this study are for patient satisfaction, acceptance of the nurses’ advice, and the use of support services for cancer.

Nevertheless, there is a lack of implementation of these approaches in Germany. OSCAR was developed to provide patients with a year of regular support in matters regarding their cancer. This is a useful addition to existing structures such as the social services provided by hospitals, which address the immediate needs of patients after hospital discharge. Improving the continuity of care and building patient competences are important for cancer patients in a fragmented healthcare system. Moreover, suitable healthcare structures and contract frameworks are necessary for delivering low-threshold access to palliative care and supporting consultations.

Experiences gathered with OSCAR could be transferred to other chronic somatic diseases which are associated with severe physical and psychological distress in patients and their relatives.

One advantage of the evaluation is the mixed-method design, which includes patient-reported outcomes, claims data from health insurance funds, and qualitative analyses of the SCNs’ project diaries. To evaluate the effects of continuous SCN support in Module 1, a prospective, non-randomized, multicenter, longitudinal study design was applied. Our patients were not assigned randomly, due to the limitation of the SCNs, who were trained and supported by the BKK insurance fund. Since randomization was not carried out, we are concerned about selection bias. Therefore, the baseline characteristics will be compared and we will control for baseline imbalance by using propensity score in our analysis. The propensity score with the inverse probability of treatment weighting (IPTW) method will be applied because the results from this method are similar to a randomized trial [23, 24]. For Module 2, we will match the intervention and control cases in order to reduce bias and confounding factors as much as possible. Moreover, given the severity of the patients’ conditions, we aimed to reduce the interview time by using the short versions of the instruments, if available, rather than applying generic instruments. In conclusion, we believe our study provides healthcare support for cancer patients. Hopefully, it can be applied for other severe chronic diseases. The evaluation of OSCAR is currently ongoing. In terms of providing healthcare support according to the patient’s needs, we expect our study to show that SCN support has a positive effect on the patient’s quality of life, health literacy, and participation in therapy planning.

Supplementary information
Supplementary information accompanies this paper at https://doi.org/10.1186/s12913-019-4585-0.

Additional file 1. ICD- and OPS-Codes. ICD-10-Codes (summary of ICD-Codes with 5 digits). OPS-Codes.

Abbreviations
API-Dm: German modified version of the Autonomy Preference Index; DCS-10: Decision Conflict Scale; DRKS: German Clinical Trials Register; EORTC: European Organisation for Research and Treatment of Cancer; GEE: Generalized estimating equation; HLS-EU-Q6: European Health Literacy Survey; IPQ: Illness Perception Questionnaire; IPTW: Inverse probability of treatment weighting; MAR: Missing at random; MICE: Multiple imputation by chained equations; OSCAR: Oncological Social Care Project; OSSS-3: Oslo Social Support Scale; PRA-D: Patient Reaction Assessment; PRO: Patient-reported outcome; SCN: Social Care Nurse; SD: Standard deviation; SN: Study Nurse

Acknowledgements
We acknowledge support from the German Research Foundation (DFG) and the Open Access Publication Fund of Charité - Universitätsmedizin Berlin.

Authors’ contributions
All authors are involved in the implementation of the study. DS and LS developed the study design. UG were involved in the statistical design of the study. JF and PG drafted the manuscript. All authors contributed to all sections of the paper. All authors reviewed the final version before submission. All authors read and approved the final manuscript.

Funding
This research project received a grant from the Innovation Fund of the Federal Joint Committee (G-BA). Reference number: 01NVF17016. The G-BA had no role in the design of the study and collection, analysis and interpretation of data and in writing the manuscript.

Availability of data and materials
Not applicable

Ethics approval and consent to participate
This project was approved by the ethics committees at Charité – Universitätsmedizin Berlin (EA2/192/17) and the Medical Association of North Rhine (2017429). The patients were enrolled in the study after providing written informed consent.

Consent for publication
Not applicable

Competing interests
The authors declare that they have no competing interests.
References

1. Bericht zum Krebsgeschehen in Deutschland 2016. Robert Koch Institut 2016. https://www.krebsdaten.de/Krebse/DE/Content/Publikationen/Krebsgeschehen/Krebsgeschehen_node.html. Accessed 02 July 2019.

2. Kunzler A, Zindel A, Znoj HU, Bargetzi M. Distress among cancer patients and their partners in the first year after diagnosis. Praxis (Bern 1994). 2010; 99(10):593–9.

3. Flattren G, Junger S, Gunkel S, Singh J, Petzold E. Traumatic and psychosocial distress in patients with acute tumors. Psychosom Med Psychol Med. 2003;53(3–4):191–201.

4. Weeks JC, Catalano PJ, Cronin A, Finkelman MD, Mack JW, Keating NL, et al. Patient, family, and partner decisional conflict in patients with metastatic non-small-cell lung cancer. N Engl J Med. 2012;367(17):1616–25.

5. Wedding U, Meran JG, Höffken K. Overtreatment in oncology: when does less become more? Ökonomologie. 2006;14(7):691–700.

6. Temel JS, Greer JA, Muzikansky A, Gallagher ER, Adamasne S, Jackson VA, et al. Early palliative care for patients with metastatic non-small-cell lung cancer. N Engl J Med. 2010;363(8):733–42.

7. Hui D, Bruera E. Integrating palliative care into the trajectory of cancer care. Nat Rev Clin Oncol. 2016;13(3):159–71.

8. Porzio R, Neugebauer S, Heckmann T, Adolf D, Kaskel P, Froster UG. Evaluation of a cancer patient navigation program ("Onkolotse") in terms of hospitalization rates, resource use and healthcare costs: rationale and design of a randomized, controlled study. BMC Health Serv Res. 2018;18(1):413.

9. Fiscella K, Whitley E, Hendren S, Raich P, Humiston S, Winters P, et al. Patient autonomy-Preferenz-Index (API-Dm). Klin Diagn Eval. 2011;4(1E):5–10.

10. Koller M, Neugebauer EAM. Methoden zur Messung von Patient-Reported Outcome (PRO). In: Pfaff H, Neugebauer EAM, Glaseke G, Schrappe M, editors. Lehrbuch Versorgungsforschung Systematik - Methodik - Anwendung. Stuttgart: Schattauer; 2017. p. 102–7.

11. Fayers PM, Aaronson NK, Bjordal K, Groenvold M, Curran D, Bottomley A, editors. EORTC Quality of Life Group. EORTC QLQ-C30 scoring manual. Brussels: European Organisation for Research and Treatment of Cancer; 2001.

12. Gaab J, Bunschoten SL, Sprott H, Ehlert U. Illness Perceptions in patients with cancer. Göttingen: Hogrefe; 2008. p. 108–117.

13. Hoebel J, Müters S, Kuntz B, Lange C, Lampert T. Measuring subjective social status in health research with a German version of the MacArthur Scale. Bundesgesundheitsbl Gesundheitsforsch Gesundheitsschutz. 2006;49(9):853–60.

14. Demographische Standards. Wiesbaden: Statistisches Bundesamt; 2004. https://www.destatis.de/DE/Methoden/Demografische-Regionale-Standards/textbaustein-demographische-standards.html. Accessed 02 July 2019.

15. O'Connor AM. User Manual - Decisional Conflict Scale. Ottawa: Ottawa Hospital Research Institute. © 1993 [Updated 2010]. https://decisionaid.ohri.ca/docs/develop/User_Manuals/UM_Decisional_Conflict.pdf. Accessed 02 July 2019.

16. Wagner EH, Ludman EJ, Aiello Bowles EJ, Penfold R, Reid RJ, Rutter CM, et al. Nurse navigators in early cancer care: a randomized, controlled trial. J Clin Oncol. 2014;32(2):259–65.

17. Bodner TE. What improves with increased missing data imputations? Struct Equ Model Multidiscip J. 2008;15(4):651–75.

18. Wagner EH, Ludman EJ, Aiello Bowles EJ, Penfold R, Reid RJ, Rutter CM, et al. Nurse navigators in early cancer care: a randomized, controlled trial. J Clin Oncol. 2014;32(1):12–8.

19. Kocalevent R-D, Berg L, Beutel ME, Hinz A, Zenger M, Härter M, et al. Social support in the general population: standardization of the Oslo social support scale (OSSS-3). BMC Psychology. 2018;6(1):31.

20. Kuss O, Blettner M, Borgemann J. Propensity score: an alternative method of analyzing treatment effects. Dtsch Arztebl Int. 2016;113(33–36):597–603.

21. Bodner TE. What improves with increased missing data imputations? Struct Equ Model Multidiscip J. 2008;15(4):651–75.

22. Wagner EH, Ludman EJ, Aiello Bowles EJ, Penfold R, Reid RJ, Rutter CM, et al. Nurse navigators in early cancer care: a randomized, controlled trial. J Clin Oncol. 2014;32(2):259–65.

23. Bodner TE. What improves with increased missing data imputations? Struct Equ Model Multidiscip J. 2008;15(4):651–75.

24. Bodner TE. What improves with increased missing data imputations? Struct Equ Model Multidiscip J. 2008;15(4):651–75.

25. Bodner TE. What improves with increased missing data imputations? Struct Equ Model Multidiscip J. 2008;15(4):651–75.

26. Wagner EH, Ludman EJ, Aiello Bowles EJ, Penfold R, Reid RJ, Rutter CM, et al. Nurse navigators in early cancer care: a randomized, controlled trial. J Clin Oncol. 2014;32(1):12–8.

27. Berezowska A, Passchier E, Bleeker E. Evaluating a professional patient navigation intervention in a supportive care setting. Support Care Cancer. 2019;27:3281–3290.