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The use of EORTC measures in daily clinical practice—
A synopsis of a newly developed manual

Lisa M. Wintner a,*, Monika Sztankay a, Neil Aaronson b, Andrew Bottomley c, Johannes M. Giesinger a, Mogens Groenvold d, Morten Aa Petersen d, Lonneke van de Poll-Franse e, Galina Velikova f, Irma Verdonck-de Leeuw g, Bernhard Holzner a on behalf of the EORTC Quality of Life Group

a Department of Psychiatry, Psychotherapy and Psychosomatics, Medical University of Innsbruck, Anichstr. 35, 6020, Innsbruck, Austria
b Division of Psychosocial Research and Epidemiology, The Netherlands Cancer Institute, Plesmanlaan 121, 1066, CX, Amsterdam, The Netherlands
c Quality of Life Department, EORTC Headquarters, Avenue E. Mounier 83, 1200, Brussels, Belgium
d The Research Unit, Department of Palliative Medicine, Bispebjerg Hospital, University of Copenhagen, Bispebjerg Bakke 23, 2400, Copenhagen, Denmark
e Netherlands Comprehensive Cancer Organisation, Gebouw Janssensenborch, Godebadkwartier 419, 3511, DT, Utrecht, The Netherlands
f Leeds Institute of Cancer and Pathology, University of Leeds, LS2 9JT, Leeds, UK
g Clinical Psychology, VU University Medical Center, De Boelelaan 1118, 1081, HZ, Amsterdam, The Netherlands

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Abstract Cancer has increasingly become a chronic condition and the routine collection of patient-reported outcomes (PROs) like quality of life is widely recommended for clinical practice. Nonetheless, the successful implementation of PROs is still a major challenge, although common barriers to and facilitators of their beneficial use are well known. To support health care professionals and other stakeholders in the implementation of the EORTC PRO measures, the EORTC Quality of Life Group provides guidance on issues considered important for their use in daily clinical practice. Herein, we present an outline of the newly developed “Manual for the use of EORTC measures in daily clinical practice”, covering the following issues: * a rationale for using EORTC measures in routine care *selection of EORTC

* Corresponding author.
E-mail addresses: lisa.wintner@tirol-kliniken.at (L.M. Wintner), monika.sztankay@tirol-kliniken.at (M. Sztankay), n.aaronson@nki.nl (N. Aaronson), andrew.bottomley@eortc.be (A. Bottomley), johannes.giesinger@i-med.ac.at (J.M. Giesinger), Mogens.Groenvold@regionh.dk (M. Groenvold), Morten.Aagaard.Petersen@regionh.dk (M.A. Petersen), lvandepoll@iaknl.nl (L. van de Poll-Franse), g.velikova@leeds.ac.uk (G. Velikova), I.M. Verdonck@vumc.nl (I. Verdonck-de Leeuw), bernhard.holzner@tirol-kliniken.at (B. Holzner).
1. Introduction

The focus of cancer care has shifted from a focus mainly on survival of patients to how patients experience their disease over the course of treatment, receive aftercare and live with cancer as a chronic condition. There is a growing demand for routine monitoring of patient-reported outcomes (PROs) like quality of life (QOL) to complement clinical data with the patient’s perspective. Although routine QOL assessments are recommended in both the literature [1–5] and evidence-based national guidelines for cancer care [6–8], they are rarely incorporated in clinical practice, though common barriers and possible solutions are well known [9–11].

As the EORTC instruments are some of the most often used PRO measures (PROMs) in oncology with high acceptability [12–14], we developed an EORTC-specific manual, to supplement the variety of available EORTC measures with guidance according to their use in routine care. Our Manual for the use of EORTC measures in daily clinical practice (available at http://groups.eortc.be/qol/) addresses all stakeholders interested in using EORTC instruments as a tool in routine patient care, providing general information on PROs and giving particular information on the EORTC measures. Corresponding to the classic manuscript style of a narrative review, the aim was to collect an extensive body of literature, extracting the most important information, evidence-based recommendations and explicitly referencing all sources of information when cited. Furthermore, considerations for an appropriate implementation strategy and issues exceeding routine care make up a large part, to give a comprehensive picture of the potential of PRO assessments.

In 2011, the International Society for Quality of Life Research developed a first version of a User’s Guide for Implementing Patient-Reported Outcomes Assessment in Clinical Practice [15], which was also published as a review article [16] and updated in 2015 [17], to provide basic information for clinicians who are interested in collecting any kind of PROs (e.g. patient-reported adverse events, functioning, health status and QOL) in their clinical practice. Following short informational paragraphs at the beginning of each chapter, the guide provides checklists in note form about needed resources, advantages, disadvantages and further reading references for each aspect dealt with. As the International Society for Quality of Life Research guide and the EORTC manual differ noticeably concerning covered issues and how information is presented, users might benefit most by combining the two sources.

This article is a synopsis of the newly developed EORTC manual, which offers a narrative review of relevant PRO literature of five different areas of interest: (1) Why EORTC PROMs should be used in routine clinical practice; (2) practical implications concerning selection, timing of assessments, scoring and presentation of data assessed with EORTC measures; (3) implementation of EORTC measures in daily clinical practice; (4) electronic data assessment and PRO-telemonitoring; and (5) further areas of application of EORTC measures and ethical considerations.

2. Why EORTC measures should be used in routine clinical practice

The presented manual uses the term ‘EORTC measures’ as an umbrella for all PROMs developed by the EORTC Quality of Life Group (QLG):

- Stand-alone questionnaires covering issues important for all cancer patients (EORTC QLQ-C30 [18], EORTC QLQ-C15-PAL [19], EORTC IN-PATSAT32 [20])
- Additional disease-specific modules [21].
- EORTC computerised adaptive testing (CAT) measures [22].
- Single items from the EORTC item library (former item bank) [23] (to create ad hoc checklists when a specific module is not available or needs to be extended)

EORTC measures undergo a rigorous development and validation procedure [24] and have been shown to be applicable across multiple countries and nationalities [25–27], as development already assesses cross-cultural
appropriateness [24], and high-quality translations [28] are available. Several studies have provided evidence that the regular use of PROMs, including the EORTC measures, facilitate and improve communication between patients and health care professionals [29–32], as clinicians’ access to PRO data makes discussion of more intimate issues more likely [33–35] and increases their awareness of patients’ functioning and well-being [33,35] without prolonging consultation time [33,36–39]. Furthermore, the use of PROMs has the potential to facilitate shared medical decision-making as patient participation is enhanced [40], to provide appropriate referral to specialists [41–44], and to enhance continuity of care [32]. Routine electronic assessment of PRO (ePRO) data enables their being linked to medical data in the electronic health records or other medical registries which support clinicians in both therapy-associated data interpretation and scientific data processing. Efforts to systematically include PROMs have increased in recent years, including in publications and white papers providing advice and best practices [45–47] and the project of the International Consortium for Health Outcomes Measurement, which is developing recommendations of common PRO data sets for different diseases, including EORTC measures [48–50].

3. Practical implications concerning selection, timing of assessments, scoring and presentation of data assessed with EORTC measures

Owing to the high-methodological quality and wide range of EORTC measures, users just need to decide what kind of data they are interested in (EORTC QLQ-C30 alone, additional disease-related modules, further issues like information or satisfaction, or EORTC CAT measures). Issues like the mode of administration (e.g. paper–pencil, electronic, phone calls) and patient burden need to be taken into account, as they should impose the least possible burden on the patients, health care professionals and clinical workflow without omitting important information.

Limited evidence-based recommendations for the timing of routine assessments with EORTC measures are available and are mainly derived from expert opinions [17,51]. In clinical practice, comprehensive knowledge of the disease and specific cancer types, treatment effects and their interaction [51], or at least elaborate hypotheses concerning this matter, should guide the timing of assessments (e.g. in advance decisions, if assessments should be time- or event-driven), as otherwise important information might be missed or patients’ scores might be misjudged [52]. In addition, a predefined rationale should guide the frequency of PRO assessments, as the evaluation of applied interventions, routine monitoring of in- or outpatients and long-term monitoring require different schedules to gather meaningful data. Few clinical models for determining assessment time-points and frequency have been developed [53,54], and further research and individual adaptation of schedules are warranted.

For EORTC measures, a scoring manual comprehensively illustrating the calculation of outcomes is available [55]. Scores might be interpreted compared to previous assessments of the same patient, related to reference populations (e.g. cancer subgroups, general population, age or gender groups) or to thresholds for clinical importance. There is a growing interest in establishing such thresholds to improve the usefulness of the EORTC QLQ-C30 in daily clinical practice [56–58], enabling the differentiation between patients with and without clinically important QOL changes, fostering its use as a screening tool and providing a reliable decision aid guiding patient-physician communication. The EORTC QLG currently funds an ongoing cross-cultural project, developing thresholds for clinical importance for all scales of the EORTC QLQ-C30.

Available literature on graphical presentation formats for the QLQ-C30 mostly focuses on group-level data [59] from clinical trials. The results of two recent studies [60,61] investigating presentation styles for PRO results from individual patients are not conclusive enough to make definite recommendations on how PRO data should be presented in clinical practice. Presumably, health care professionals and patients may differ in their preferences regarding types of graphical presentation and the amount of information included.

Specific training and information materials tailored to patients’ or health care professionals’ needs should be used to enable them to correctly understand the presented results of the EORTC measures. Fig. 1 provides examples of presentation styles used by members of the EORTC QLG.

4. Implementation of EORTC measures in daily clinical practice

Integration of PRO assessments using EORTC measures into clinical routine is a complex health care intervention with multiple interacting components on different levels of the clinical system, all of which are sensitive to multiple influences and barriers [62,63]. Based on available recommendations on PRO implementation into clinical practice [e.g. Refs. [10,17,64–67] and its own experience, the EORTC QLG proposes some practical approaches to support successful integration (please refer to Table 1).

5. Electronic data assessment and telemonitoring

Routine ePRO assessment with the EORTC measures provides real-time results, can reduce time and labour...
needed for data entry and calculation as well as continuous expenditures (e.g. for printing, though costs for software support might cancel out these savings) provides high-data quality and improved care may be achieved. Paper—pencil versions can serve as a back-up or assessment method for those patients who would otherwise decline assessment (e.g. older people [78,79]), as studies report a general equivalence between paper—pencil and computer-based versions of PROMs [80]. At present, jointly with the EORTC QLG, Mapi [81] is developing guidelines for graphical presentation formats of electronic versions of the EORTC measures.

Infrastructural requirements for ePRO comprise both technical and educational aspects. A comprehensive IT infrastructure is needed and includes technical devices for data collection and output (workstations, hand-held devices, preferably equipped with touchscreens), appropriate software solutions and network facilities for data transmission, storage and back-up, technical support and updates. Furthermore, user skills regarding computer literacy and open-mindedness towards PRO assessments are of special importance.

There are a variety of systems available which allow ePRO [82–84], but systems vary remarkably concerning their features. In choosing software for ePRO of EORTC measures, it should be carefully checked if it is flexible enough and able to provide all desired functionalities and possibly any future necessary adoptions. Since 2009, the EORTC QLG has been supporting the development of the CHES.EORTC software [85], which allows ePRO in daily clinical practice, clinical studies and long-term follow-up via telemonitoring and provides a data exchange interface for clinical information systems based on the Health Level Seven standard as well as an elaborated graphical results presentation of patient-level data [86]. Using CHES.EORTC within a patient portal accompanied by educational and self-help material is possible as well.

Continuous assessment of patients’ health status and QOL outside the hospital via telemonitoring can add to patient care. Especially patients undergoing active outpatient treatment might benefit from telemonitoring, as side effects and treatment-related symptom burden might occur outside the hospital [87]. Alert systems
Table 1

<table>
<thead>
<tr>
<th>Understand current practice before applying strategies for integration.</th>
<th>Existing patient care pathways, knowledge and attitudes towards EORTC measures as well as the skills of potential adopters should be assessed and resources and barriers analysed [68]. A baseline assessment could identify elements relevant for the adaption of practice change, such as benefits, harms and costs [69].</th>
</tr>
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<tr>
<td>Engage all relevant stakeholders in all stages of implementation.</td>
<td>This should include the introduction of existing national guidelines, providing the opportunity for health care professionals to discuss current practice [70], engaging senior clinicians, nurse champions and opinion leaders and assigning a coordinator throughout the implementation process [16].</td>
</tr>
<tr>
<td>Collaborate to define the goals and expectations for routine assessments with EORTC measures up front to ensure relevance.</td>
<td>Health care professionals’ active participation in the ongoing integration process ensures that they regard assessments with EORTC measures to be relevant for their clinical practice, thereby facilitating uptake of the new intervention into clinical routine.</td>
</tr>
<tr>
<td>Make data actionable.</td>
<td>Automatic ‘flagging’ of clinically important scores, providing longitudinal interpretation of what signifies a clinically important change in EORTC measures data as well as linking of results with clinical practice guidelines and interventions [30,65,70] are examples how EORTC measures data can be made available for health care professionals.</td>
</tr>
<tr>
<td>Provide training, coaching and support for health care professionals as well as patients and their informal caregivers.</td>
<td>A multifaceted, interactive approach [71–73] is expected to be more effective for practice change than large-scale, passive education materials [74] and detailed reports on such specific programs are available [75,76]. At the patient level, disease, cultural and personal aspects should be considered to reduce response burden [70].</td>
</tr>
<tr>
<td>Evaluate integration process and outcome.</td>
<td>Defining outcome as well as process indicators together with health care professionals will help them reflect on the process of integration, to identify obstacles and to adapt implementation strategies where necessary [30]. Regular reviews will determine progress, elicit problems and engage health care professionals in identifying efficient strategies to overcome barriers [68].</td>
</tr>
<tr>
<td>Consider organisational context</td>
<td>Although implementation is not recommended concurrent with other major organisational changes [75], organisational adoption will be necessary to support sustainability of routine assessment of EORTC measures and their being embedded in integrated cancer care pathways.</td>
</tr>
<tr>
<td>Long-term evaluation of effective integration.</td>
<td>Long-term follow-up is necessary to determine if short-term changes persist and whether surrogate outcomes point towards benefits [63]. Quasi-experimental, observational studies or service development and evaluation models based on quality improvement and implementation science methods might be more informative than randomised controlled trials [77].</td>
</tr>
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enable health care professionals to recognise occurring symptoms in real time despite the geographic distance between the patient and medical services, contact patients and intervene appropriately [88]. Research can also benefit from telemonitoring with EORTC measures as such data can be merged with routine assessments at the hospital and knowledge of the disease itself and treatment-related symptoms might be deepened.

Telemonitoring seems to be feasible for a considerable percentage of cancer patients [89,90] and patients show more and more interest in the use of electronic portals [91], which could provide, next to trustworthy educational and illustrating material, the possibility to complete EORTC measures as well. There is some evidence suggesting several benefits of using patient portals including the feeling of being better prepared for clinical appointments, development of better coping strategies, higher satisfaction with treatment choices, and better adherence to medical advice [92].

Nonetheless, efforts need to be made to include patients who are less likely to engage with electronic assessments due to unfamiliarity with or missing access to computer technology, lower interest in electronic health activities [93,94], or special needs (but auxiliary tools like adjustable font sizes, voice output or interactive voice response, additional input devices, telephone interviews may overcome this problem). Furthermore, implementation of telemonitoring needs to include a clear strategy on how to proceed if patient data indicates problems needing acute medical attention and how patients are effectively educated about the complementary nature of EORTC measures. Patients need to be aware that telemonitoring and using provided educational material and self-help advice must not replace or delay their visit to the hospital or doctor if they are feeling unwell.

6. Further areas of application of EORTC measures and ethical considerations

Routinely collected EORTC measures data can be a valuable source for comparing outcomes with predefined benchmarks, evaluation of practices, care pathways, policies, treatment technologies and disclosing systematic differences in distribution of health care among patients [95–97]. PRO data might even be used for certification
requirements, as they stand for both the incorporation of patients’ perceptions and patient empowerment.

Even though quality assurance can be a promising area for the use of PRO, there is a strong need for further research, as the interests of different stakeholders vary widely. So far, there are no standards on how PRO data have to be collected to consider the needs of all stakeholders without privileging or disadvantaging any of the parties involved, although some efforts have been undertaken to develop recommendations concerning this matter [98].

In the near future, EORTC measures can also be used for cost-utility analysis, and therefore for comparative effectiveness research [99]. This has become possible with a recent project of the EORTC QLG and the international MAUCa (Multi-Attribute Utility in Cancer) group, within which the EORTC QLU-C10D measure was developed, a utility instrument enabling the estimation of quality adjusted life years using the EORTC QLQ-C30 data [100]. Such analyses may focus on new treatment modalities or compare existing options, setting their monetary costs and QOL benefits and impairments against each other. Value sets have currently been developed for Australia, Germany, Poland, France and Italy and valuations in further European countries are planned.

Regarding ethical issues, it is important to balance the effort PRO assessments impose on patients with the expected benefit. Which and how many EORTC measures are used for data collection needs to be carefully chosen without underestimating the patients’ ability and willingness to complete EORTC measures because they seem to be ‘too ill’. The systematic exclusion of very ill patients alters data at a group level and may deprive exactly those patients who are in special need of particular interventions (e.g. referral to psychosocial services).

Before patients start completing EORTC measures, they need to be extensively informed about the purpose of assessments, issues concerning data confidentiality and security and their rights to withdraw from assessments and request destruction of data. Furthermore, disclosure needs to address common myths about electronic health interventions and telemonitoring (e.g. the fear that such interventions are offered because of disease progression, that the patient’s own coping skills are insufficient [101], that there will be loss of personal contact with health care team).

Privacy is not only an issue if ePRO is conducted (adequate IT structure to ensure maximum data protection for collection, transmission and storage) but is also related to the environment of data assessment, as patients should be able to complete EORTC measures unobserved on their own in a private area, without any time constrains and the possibility to get help if necessary.

Protecting the confidentiality of patients’ data might interfere with the usability of data in clinical routine.

Some patients might not want all members of their clinical team to see their data, because they report sensitive information (e.g. sexual issues) or might fear negative consequences (e.g. following a report of low satisfaction with care). It is still a challenge to find adequate ways for data processing and access levels to preserve the interests of the parties involved.

7. Conclusion

This article briefly summarises the issues covered by our newly developed Manual for the use of EORTC measures in daily clinical practice, which aims at fostering the successful implementation of regular PRO assessments in routine patient care. The field of PRO assessments is continuously evolving and issues change regarding their importance and relevance. Medical, technical and regulatory advances and new scientific insights need to be taken into account when recommendations about the routine use of PROs are given. Consequently, guidelines like the presented EORTC manual need to be periodically updated. Possible topics to be included in a future version of the manual could be a more in-depth discussion on the use of EORTC measures for continuity of care, a critical appraisal of the possible use of the EORTC measures for quality assurance, a summary of the Mapi/EORTC QLG guidelines on the graphical presentation of EORTC measures in ePRO software, and the possible linkage between EORTC measures and self-management programs. As this manual is made for end-users of EORTC measures, any comments and suggestions for improvement and further development of the manual are warmly welcome: http://groups.eortc.be/qol/contact-us.

Conflict of interest statement

Bernhard Holzner is owner of the intellectual property rights of the CHES software. The other authors declare no conflicts of interest.

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