Patient-reported outcomes in clinical trials

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Abstract
Patient-reported outcomes (PROs) are progressively being included in clinical trials to provide information about treatment benefits identified by the patients themselves that extend the data on traditional clinical trial endpoints, such as disease free survival, overall survival, progression-free survival, and response rate. PROs may have a greater impact for patients than other endpoints. For example, patients may be prepared to forgo some increase in progression-free survival in return for reduced treatment-related toxicity. PROs may also be an indicator of disease response and have value as prognostic factors. This article discusses the way PROs can be defined and incorporated into clinical trials to enhance the value of clinical trials data and improve the understanding of the clinical benefits of a specific treatment, not only for health care professionals, but for patients and caregivers. The importance and relevance of a patient-centered perspective and shared decision making in defining value and determining treatment benefit is increasingly recognized. However, despite the acknowledged value of PROs, their inclusion in clinical trials remains far from ideal. New guidelines from the research community and technological improvements in data collection and analytics will increase the quality and the importance of PROs as standard methods for the evaluation of medical studies and in the drugs approval process.

Key words: clinical benefit, clinical trial endpoints, CONSORT PRO extension, patient-reported outcomes, reporting quality

Introduction
The most frequent endpoints of clinical trials are: disease free survival (DFS) and overall survival (OS), frequently used in the adjuvant or curative setting; progression-free survival (PFS), OS and response rate (RR), more often employed in the metastatic diseases. Nevertheless, these parameters are not completely able to reproduce the clinical benefits and risks of the therapies. Since the last decade, patient-reported outcomes (PROs) have been progressively included in clinical trials, in order to produce more complete information about the benefits of treatments as reported by the patients themselves. PROs could have a more significant impact for patients than other endpoints, as shown by Havrilesky et al. in a pilot study, which reported that women with ovarian cancer are ready to sacrifice some of meaningful endpoints, such as PFS time, for less heavy treatment-related toxicity [1]. Moreover, improvements in PROs may be correlated with disease response, as suggested in a meta-analysis conducted in 2006 [2]. In some studies, PROs have been recognized as prognostic factors [3, 4]. The association of quality of life (QoL) parameters and symptoms (e.g. pain and weight loss) with clinical characteristics and socio-demographic variables (e.g. age, gender, performance status, stage of the disease) can increase the prediction of OS by 6%, in comparison with the use of clinical characteristics and socio-demographic variables alone [3]. In a recent study presented by Roncolato et al., baseline QoL data reported that low physical function, role function, global health status and high abdominal or gastrointestinal symptoms are predictors for OS and for stopping chemotherapy early in platinum-resistant/refractory ovarian cancer [5].

Definitions and measures
PROs are defined as any report of the status of a patient’s health condition, originated from the patient himself, without interpretation of the patient’s response by the clinician or anyone else [6]. PROs can include different aspects
of the health condition, such as QoL, health related QoL (HRQoL), patient’s satisfaction with care, symptoms, pain, psychological distress, self experience and patient-reported adherence to therapy. These data are collected in most of cases by the patients directly through standardized and validated questionnaires, such as the European Organization for Research and Treatment of Cancer (EORTC) QoL questionnaires or Functional Assessment of Cancer Therapy (FACT) questionnaires. A possible alternative is to collect the spontaneous feedback of patients, with the help of caregivers, especially in the elderly population and/or in the palliative setting, when patients are not independent [6].

Others measures have been developed to assess the value of therapies, including the widely used metric of quality-adjusted life-years (QALYs). QALYs are an indicator of disease burden, that includes both quality and quantity of life lived. The QALY’s measure, collected on a selected population of patients, provides an indication about the benefits obtained from medical procedures in terms of QoL and survival. QALYs is used to evaluate and compare the effectiveness of different treatments, enabling the choice of the most cost-effective one. QALYs can be used as parameter for decision making for specific patients, but acquires more importance when is used for statistical and economical evaluations [7, 8]. Since published studies have demonstrated an under-reporting of patients’ symptoms and their agreement between the evaluations of patients and clinicians, the Patient Reporting of Common Terminology Criteria for Adverse Events (PRO-CTCAE) library has been developed [6, 9, 10]. It includes 78 symptomatic side effects and adverse events (AEs) commonly reported in oncology clinical trials and assesses their frequency, severity and interference with daily activities. The patients’ AEs reports could give additional information particularly in the target therapy era, when drugs should be taken for an extended time, in order to more deeply analyze the overall benefit [10].

Composite endpoints including PROs, tumor response and survival outcomes have been developed in order to enhance the value of clinical trial results and to assist the deeper understanding of the entire clinical benefits of a specific treatment, both by health care professionals (HCP), patients and caregivers [4]. Different composite endpoints for PROs have been proposed, such as clinical benefit response (CBR) endpoint, time until definitive deterioration (TUDD) and overall treatment utility (OTU) [4, 11-13]. In this context, CBR endpoint includes pain (result of both analgesic use and patient-reported pain intensity scale), clinician-recorded performance status and weight. CBR was initially defined by Burris et al. and, subsequently, widely used across the literature [11]. CBR should be not confused with the definition of clinical benefit reported in tumor-centered trials, where it indicates the changes of tumor size after treatments (complete response, partial response and stable disease). TUDD is calculated in different ways according to the studies in which it is used. Bonnetain et al. evaluated QoL every 8 weeks until death through these parameters: global health, emotional functioning, physical functioning, fatigue and pain. Bonnetain’s TUDD was calculated as the elapsed time between the start of the study and the first definitive decrease in QoL parameters (mentioned above) by 5 or more points [12]. OTU incorporates tumor response, serious adverse events, grade >3 non-hematological toxicity and death [13]. However, there is not a consensus about the treatment related symptoms in composite endpoints, which are not yet widely validated and used.

The relevance of patient centeredness and of patient perspective in defining value has been recognized by the American Society of Clinical Oncology (ASCO) and the European Society for Medical Oncology (ESMO). In order to help HCP and patients to share decision making, ASCO developed a value framework, divided into two different versions, one for curative settings and one for advanced disease. Both models consider clinical benefits (including the most frequently used endpoints, OS, PFS and RR) and toxicity (this value represents the delta between the toxicity of the new agent and the comparator therapy). For advanced disease, the model includes a PROs endpoint, as “bonus point”, recognizing improvements in cancer-related symptoms and prolongation of the treatment-free interval. The prolongation of treatment-free interval is supposed to be a good-health substitute, since it indicates a vacation from drug toxicities [7].

ESMO has also recently developed a Medical Oncology Magnitude of Clinical Benefit Scale (ESMO-MCBS), in order to evaluate the meaningful benefits obtained from a new treatment. The ESMO-MCBS incorporates OS, PFS and QoL. In particular, QoL can upgrade or downgrade the evaluation of the new drug, respectively when a statistically significant improvement in QoL or its delayed deterioration have been recorded or, on the contrary, when an absence of QoL advantage has been reported [14].

**How to include PROs in clinical trials: the CONSORT PRO extension**

Although PROs are important endpoints in clinical decision making, and even if PRO measures are included in studies, only few randomized controlled trials (RCTs) adequately report PRO data. In a review published in 2011, Brandugue et al. reported that in an identified sample of 794 RCTs,
PROs were primary endpoints in just 26% of the total trials; in the 56% of the total trials where the rationale for the selected PROs was documented, 28% of the RTCs gave information about missing data and 64% of the RTCs analyzed the PRO results in the context of other endpoints [15]. The sub-optimal protocols for reporting PROs in clinical trials are also confirmed by two recent articles focused, respectively, on ovarian cancer and advanced breast cancer, in particular regarding timing of administration of PROs instruments, monitoring of PROs compliance, handling of missing data, the analysis plan for PROs and results discussion [16, 17].

Since the interpretation of PROs data requires accurate and standardized reporting, the Consolidated Standards of Reporting Trials (CONSORT) Statement developed a PROs centered extension. The CONSORT PRO guidance identified 5 objectives that must be present in all RCTs in which PROs are primary or significant secondary endpoints:

1. PROs should be recognized as primary or secondary endpoints in the abstract;
2. if a multidimensional PROs instrument is used, the hypothesis and relevant fields must be described;
3. the evidence of the PROs tool’s validity (evaluate what they are designed to measure) and reliability (get the same answer repeatedly) should be explained or mentioned;
4. the statistical approaches to manage the missing data should be clearly explained, since missing data reduce power of the study and is a potential cause of bias;
5. PRO-specific restrictions of study results and generalizability of findings to other populations and clinical practice should be discussed.

In the results section, baseline PROs data must be presented, since they could be useful to assess the importance of trial findings. If possible, additional PROs or the elements of composite PRO scores should be analyzed in the main publication (or in a companion second publication), in order to reduce selection bias of reporting only significant results. In fact, some studies do not take into account the clinical relevance of PROs, especially in RCTs where there are discrepancies between PROs themselves and survival endpoints [18].

When and how PROs could be included in clinical trials

In advanced disease, the objective is to improve survival without degradation of QoL. For this reason, the inclusion of PROs assumes a particular relevance, more particularly when the first goal is to improve palliation of symptoms. Symptoms control or improvement, with the consequent advantage in HRQoL, represents a more important and substantial clinical benefit than RR (according to Response Evaluation Criteria In Solid Tumors [RECIST] criteria) or PFS alone. Collection of HRQoL is important not only during the treatment, but also after and during subsequent lines. Data might evidence a delay in the onset of cancer-related symptoms and a postponement of following treatments. The extension of HRQoL data recording after the end of therapy could evidence late onset toxicities and related symptoms. In advanced disease, even if the treatment is becoming similar to other chronic diseases like hypertension or epilepsy, the approval of new drugs is exceptionally based only on reduction of symptoms [19]. Even if the Food and Drug Administration (FDA)’s Guidance to Industry in 2007 suggested that symptom endpoints can be sufficient in the regular drugs approval process, in practice only mitoxantrone in metastatic prostate cancer has been approved based on symptoms palliation [20, 21].

A patient-centered outcome research, based on standardized PROs, can also add important value in adjuvant therapy and in asymptomatic relapse. In the adjuvant setting, assessing acute and long term side-effects of the treatment is an important goal. In some cases, for a marginally improvement in survival, these patients could decide to refuse the proposed treatment or to postpone it at the onset of symptoms. In this setting the collection of HRQoL data with reference to late onset toxicities acquires more importance, such as anthracycline related heart impairment and peripheral neuropathy, that could significantly worsen daily activities.

In the area of targeted therapies and immune checkpoint inhibitors prescribed for long periods and sometimes in the maintenance setting, the evaluation of PROs, in particular the ones related to chronic toxicities and to adherence of therapies, will continue to become more important. Many of the targeted therapies are small molecules with an oral administration and the patient’s compliance assumes a fundamental role for their efficacy [6]. The new agents have different toxicity profile in comparison to chemotherapy and the value of using RECIST criteria to assess response is under review. In this setting, inclusion of PROs data as endpoints can help in understanding trial results and in evaluating the overall benefit. For example, the proportion of patients that reached a significant improvement in symptoms from baseline at a predefined time point is perhaps more important than a modest rate of disease progression based on RECIST criteria or a marginal PFS difference of some weeks [22].

Future perspectives and conclusion

The importance of including PROs in clinical trials is largely recognized today, according to the CONSORT PRO ex-
tension recommendations. Nevertheless, a recent review by Bylicki et al. found that 62% of RCTs did not report PROs, while the other ones lacked information about items associated with methods of PROs collection and analysis (just 16% of the RCTs recorded methods for data collection and only 37% described the management of missing data) [23]. The presence of a dedicated secondary manuscript was an independent predictor of overall quality score [23]. Starting from 2007, the FDA’s Guidance to Industries included symptoms as important endpoint for regular drug approval. For this reason, it is important to encourage the inclusion of PROs in the endpoints of RCTs. The new technologies, such as the digital conversion of the traditional paper questionnaires and the big data analytics, will increase the capture and statistical analyses of PROs data, with a more strict observation of the pre-scheduled detection and a reduction of the amount of missing data. A positive trend for reduction of missing data and an increment in compliance rate (between 84.7 and 97.2%) have been reported recently in Alliance and Mayo Clinic trials [24]. The research community guidelines, developed during the past years based on several different studies, and the technological improvements about data collection and analytics will increase the quality and the importance of PROs as standard methods for the evaluation of medical studies and in the approval processes for new drugs.

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Conflicts of Interest
The Authors declare there are no conflicts of interest in relation to this article.

References


