Original Research

Self-evaluation of duration of adjuvant chemotherapy side effects in breast cancer patients: A prospective study

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Abstract

Background: We recently reported that self-evaluation of the incidence and severity of treatment-related side effects (TSEs) using a National Cancer Institute (NCI) Common Terminology Criteria for Adverse Events (CTCAE) v4.0-based questionnaire was feasible and more informative than doctor reports in patients undergoing standard adjuvant chemotherapy for operable breast cancer. Here, we compare self- and doctor-evaluated day of onset and duration of TSEs in the same population.

Patients and methods: Six hundred and four patients were enrolled at 11 sites in Italy. CTCAE v4.0 definitions of grade of severity of nausea, vomiting, constipation, anorexia, dysgeusia, diarrhea, fatigue, pain, paresthesia, and dyspnea were translated into Italian and rephrased. Questionnaires were administered after the first and third
The collection and analysis of treatment-related side effects (TSEs) are critical to the management of patients with cancer both in clinical trials and in daily practice. Systems like the National Cancer Institute (NCI) Common Terminology Criteria for Adverse Events (CTCAE) are integral to clinical trials where measures of treatment-related toxicity are critical for establishing the risk-benefit ratio of new treatments. Interestingly, a number of papers have reported moderate to substantial underestimation of TSE reporting when investigator-collected data are compared to corresponding data provided by patients using adapted questionnaires. For this reason, the NCI has promoted the development of a “patient” version of the CTCAE (patient-reported outcome (PRO)-CTCAE), which is currently being translated and validated in different languages and integrated, together with more established PROs, in clinical trials.

If patient- and doctor-reported TSEs are discrepant in the prospective clinical trial context, it is likely that TSE underestimates are even greater in patients receiving standard treatments in clinical practice, as there is no standardized system to collect them. TSE underreporting may have consequences for quality of life and disease-related outcomes. For this reason, the NCI has promoted the development of a “patient” version of the CTCAE (patient-reported outcome (PRO)-CTCAE), which is currently being translated and validated in different languages and integrated, together with more established PROs, in clinical trials.

Therefore, we developed a CTCAE v4.0-based patient questionnaire by translating grade of toxicity definitions into Italian and administering it to a large cohort of patients with breast cancer receiving adjuvant chemotherapy at different Italian institutions. We showed that doctors underestimate TSE incidence and severity in clinical practice, with a more pronounced effect in high-volume centers. However, our questionnaire also reported the day of onset and duration for each TSE considered, allowing us to here to analyze and compare these two patient- and doctor-reported TSE dimensions.

2 | PATIENTS AND METHODS

The details of our prospective trial evaluating a CTCAE v4.0-based patient questionnaire are reported elsewhere. Our ten-item paper questionnaire included nausea, vomiting, constipation, anorexia, dysgeusia, diarrhea, fatigue, pain (generic), paresthesia, and dyspnea. For each item, definitions and severity grades were translated into Italian from the CTCAE version 4.0. In addition to severity grade, the questionnaire contained fields to record the day of onset for each item (where the day of chemotherapy administration was day one), the duration in days, and persistence at the time of questionnaire administration (Figure S1). When the Italian translation of CTCAE version 4.0 became available and was endorsed by the Italian Association of Medical Oncology in mid-2011, the questionnaire was rechecked and there were no translational discrepancies.

Patients were instructed to complete the questionnaire at the end of the first cycle (usually on day one of the second cycle of planned chemotherapy) and at the end of the third cycle (usually on day one of the fourth cycle of planned chemotherapy). Dedicated nurses provided instructions on how to complete the questionnaires at the time of obtaining informed consent, with no further assistance given during the study. Patients were also provided with a diary to help record
onset and duration of specific adverse events, but this was not part of the formal study materials. At each participating Institution, dedicated nurses extracted side effect information from the medical records of patients and filled in “doctor” questionnaires at matching time-points. These questionnaires were collected exclusively by nurses and were not available to the treating doctors.

2.1 | Statistical analysis

Medians and means are reported together with their ranges and standard errors, respectively. Paired medians and means were compared by the Wilcoxon signed-rank test and by Student’s $t$ test for paired data. Proportions in unrelated samples were compared by the chi-squared test. Statistical significance was set at $P < 0.05$ (two-tailed).

3 | RESULTS

Of 604 patients with early breast cancer enrolled in the study at 11 Italian Institutions, three withdrew informed consent before the first cycle of chemotherapy. Patient demographics are summarized in Table 1. Overall, 596 and 581 patient questionnaires were collected after cycles one and three of adjuvant chemotherapy, respectively. Of these, 594 and 573 had a corresponding (doctor) questionnaire extracted from the medical charts at the same time-points, respectively. In patients reporting TSEs, the average proportions of questionnaires with complete information on day of onset and duration were 70% and 95%, respectively (first questionnaire), and 75% and 95%, respectively (second questionnaire, Tables S1 and S2). Corresponding figures for doctor reports were 29% and 95%, respectively (first questionnaire), and 31% and 97%, respectively (second questionnaire, Tables S1 and S2).

The median day of onset and median durations as reported by patients and described by doctors in the first and second questionnaires are summarized in Tables S3 and S4. As data on day of onset were missing in a high proportion of doctor questionnaires, patient and doctor reports of day of onset could not be formally compared.

The median duration values of each of the ten adverse events as reported by patients and as described by doctors after cycles one and three were only compared for pairs of charts where both patients and doctors reported each item as present (any grade). After cycle one, with the exception of vomiting, pain, neuropathy, and dyspnea, the median duration of symptoms was longer when reported by patients than doctors, with notable differences for nausea (4 vs 2 days; $P < 0.01$), constipation (3 vs 1 days; $P < 0.01$), anorexia (3 vs 1 days; $P < 0.01$), diarrhea (3.5 vs 1 day; $P = 0.02$), and fatigue (4 vs 1 days; $P < 0.01$) (Table 2). Results were similar after the third cycle, with the difference that vomiting also lasted longer when reported by patients than doctors and that the difference in the duration of diarrhea was not statistically significant (Table 3).

To analyze the combined effect of differences in reporting incidence and duration, the mean percentages of cycle days with a certain side effect were calculated for each TSE (total days with the side-effect/total days from the date of chemotherapy administration and that of data collection*100) as reported by patients and doctors. Here, “0” days described a TSE that either did not occur or that was missed by the patient or the doctor. Results for the first questionnaire are displayed in Figure 1 and in Table S5; the mean percentage of cycle days with each TSE was significantly higher when reported by patients compared to doctors. Similar results were obtained with the second questionnaire (data not shown).

4 | DISCUSSION

Here, we exploited our CTCAE V4.0-based questionnaire study to integrate data on time of onset and duration of ten common TSEs experienced by breast cancer patients treated with adjuvant chemotherapy in clinical practice.
In our prior publication comparing patient- and doctor-reported TSE incidence and severity, we described a high rate of questionnaire completion, except for the field indicating persistence of the TSEs at the time of the visit. When reviewing the dataset, despite a high completion rate, a substantial number of “day of onset” entries in both the patient and doctor questionnaires were “0” values. As the day of onset had to be counted from the day of the last chemotherapy administration (day one), we considered all the 0 values as incorrect and treated them as missing values. Consequently, for this toxicity dimension, we were unable to perform a formal statistical comparison of paired patient and doctor questionnaires for all the items because of insufficient paired data. The descriptive analysis reported in Tables S3 and S4 suggested no major differences in the median day of onset but a larger variability in patient-reported nausea, vomiting, constipation, anorexia, and dysgeusia.

Conversely, data on TSE duration were of better quality and reported in over 90% of patient and doctor questionnaires. In paired comparisons, patients reported longer duration after the first and third cycles for six and seven of the ten items, respectively. By combining information on incidence and duration, we determined that the mean percentage of cycle days with a certain toxicity was significantly higher in patient than doctor reports. These results indicate major discrepancies in the subjective experience of patients given the opportunity to self-collect TSE to medical chart reports during routine visits at oncology centers compared to doctor reports. Nevertheless, our data suggest that longitudinal TSE evaluations by patient self-administered questionnaires are

<table>
<thead>
<tr>
<th>Item</th>
<th>N paired questionnaires reporting the item</th>
<th>Duration patient median (range)</th>
<th>Duration doctor median (range)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nausea</td>
<td>175</td>
<td>4 (1-21)</td>
<td>2 (1-21)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Vomiting</td>
<td>54</td>
<td>2 (1-6)</td>
<td>1 (1-10)</td>
<td>N.S.</td>
</tr>
<tr>
<td>Constipation</td>
<td>52</td>
<td>3 (1-20)</td>
<td>1 (1-14)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Anorexia</td>
<td>34</td>
<td>3 (1-15)</td>
<td>1 (1-10)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Dysgeusia</td>
<td>38</td>
<td>2 (1-21)</td>
<td>1 (1-20)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>17</td>
<td>3.5 (1-13)</td>
<td>1 (1-7)</td>
<td>0.02</td>
</tr>
<tr>
<td>Fatigue</td>
<td>112</td>
<td>4 (1-21)</td>
<td>1 (1-14)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Pain</td>
<td>28</td>
<td>1 (1-15)</td>
<td>1 (1-15)</td>
<td>NS</td>
</tr>
<tr>
<td>Paresthesia</td>
<td>13</td>
<td>1 (1-6)</td>
<td>1 (1-6)</td>
<td>NS</td>
</tr>
<tr>
<td>Dyspnea</td>
<td>11</td>
<td>1 (1-6)</td>
<td>1 (1-6)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Number in parentheses indicate paired patient and doctor questionnaires available for the analysis of incidence. NS, nonsignificant.

<table>
<thead>
<tr>
<th>Item</th>
<th>N paired questionnaires reporting the item (grade ≥1)</th>
<th>Duration patient median (range)</th>
<th>Duration doctor median (range)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nausea</td>
<td>177</td>
<td>4 (1-18)</td>
<td>2 (1-21)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Vomiting</td>
<td>53</td>
<td>2 (1-8)</td>
<td>1 (1-5)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Constipation</td>
<td>54</td>
<td>3 (1-20)</td>
<td>1 (1-18)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Anorexia</td>
<td>35</td>
<td>4 (1-15)</td>
<td>1 (1-15)</td>
<td>0.01</td>
</tr>
<tr>
<td>Dysgeusia</td>
<td>48</td>
<td>2 (1-21)</td>
<td>1 (1-21)</td>
<td>0.03</td>
</tr>
<tr>
<td>Diarrhea</td>
<td>12</td>
<td>3 (1-12)</td>
<td>1 (1-7)</td>
<td>0.06</td>
</tr>
<tr>
<td>Fatigue</td>
<td>103</td>
<td>4 (1-40)</td>
<td>1 (1-21)</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Pain</td>
<td>27</td>
<td>2 (1-21)</td>
<td>1 (1-21)</td>
<td>NS</td>
</tr>
<tr>
<td>Paresthesia</td>
<td>16</td>
<td>1 (1-10)</td>
<td>1 (1-10)</td>
<td>NS</td>
</tr>
<tr>
<td>Dyspnea</td>
<td>22</td>
<td>1 (1-19)</td>
<td>1 (1-18)</td>
<td>NS</td>
</tr>
</tbody>
</table>

*Numbers indicate the pairs of questionnaires where both the patient and the doctor indicated the occurrence of the side effect and for which information on duration was present.
feasible and could improve the management of patients undergoing adjuvant chemotherapy for breast cancer.\textsuperscript{18,19}

The metrics and tools available to longitudinally evaluate side effects to dissect and exploit the different TSE dimensions in clinical trials evaluating newer treatments have recently been reviewed and discussed.\textsuperscript{20,21} This issue is becoming particularly relevant to new treatment paradigms such as the chronic administration of low-dose chemotherapy or use of biologicals. Longitudinal assessment requires several TSE dimensions to be collected during treatment to feed databases that can generate appropriate metrics. In this respect, a tool like our questionnaire or the more standardized PRO-CTCAE with integrated day of onset and duration could provide such data. Although emphasis is often given to acute toxicities from cytotoxic chemotherapy, longitudinal assessment of the dimensions of timing of onset and duration may improve TSE management in routine clinical practice. Questionnaires like our own could also be converted into Web-based tools or smartphone “eHealth” applications for the patient to complete at home, with remote access by medical staff. Some nonrandomized studies of electronic TSE data capture have been conducted, highlighting the promising potential of this approach.\textsuperscript{22-24} Importantly, a recently published randomized trial showed that advanced cancer patients undergoing treatment managed with a system integrating a Web-based version of the PRO-CTCAE and remote control with appropriate actions when needed (ie, patients reporting significant worsening of a certain TSE) had, compared with usual care, improved overall survival.\textsuperscript{15}

In conclusion, here we confirm that patients with breast cancer undergoing adjuvant chemotherapy can self-assess and report the day of TSE onset and describe their duration. The duration of most TSEs was longer when reported by patients rather than doctors, and the combined effect of longer duration and underreporting was an increase in the percentage of cycle days with all TSEs. We are now implementing an electronic version of our questionnaire in the routine management of patients with breast cancer undergoing adjuvant chemotherapy at our institution.

CONFLICT OF INTEREST

Speaker’s honoraria: Astra Zeneca, Novartis, Roche; Travel grants: Astra Zeneca, Roche; Consulting or Advisory Role: Bristol-Myers Squibb, Merck, Roche; Travel, Accommodations, Expenses: Bristol-Myers Squibb; All remaining authors have declared no conflict of interest.

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REFERENCES


**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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