Patient-reported outcomes in patients with a germline BRCA mutation and HER2-negative metastatic breast cancer receiving olaparib versus chemotherapy in the OlympiAD trial

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Abstract  Background: The phase III OlympiAD study (NCT02000622) showed a statistically significant progression-free survival benefit with olaparib versus chemotherapy treatment of physician’s choice (TPC) in patients with a germline BRCA mutation and human epidermal
Health-related quality of life; EORTC QLQ-C30; Breast cancer; BRCA growth factor receptor 2-negative metastatic breast cancer. From this study, we report the effect of olaparib on health-related quality of life (HRQoL).

Methods: Patients were randomised 2:1 to olaparib monotherapy (300 mg twice daily) or single-agent TPC. The primary HRQoL end-point was mean change from baseline in the two-item global health status/QoL score determined from patient-completed European Organisation for Research and Treatment of Cancer Quality of Life Questionnaire Core 30-item module (EORTC QLQ-C30) questionnaires and assessed using a mixed model for repeated measures. Symptoms and functioning domains, best overall response and time to deterioration of QoL were also evaluated.

Results: Overall questionnaire compliance rates were 93.2% for olaparib and 76.3% for TPC. Between-treatment global health status/QoL comparison showed a significant improvement in the olaparib arm versus the TPC arm, with mean change of 3.9 (standard deviation 1.2) versus −3.6 (2.2), a difference of 7.5 points (95% confidence interval [CI]: 2.48, 12.44; P = 0.0035). A higher proportion of patients in the olaparib arm showed a best overall response of ‘improvement’ in global health status/QoL (33.7% vs 13.4%). Median time to global health status/QoL deterioration was not reached in olaparib patients and was 15.3 months for TPC patients (hazard ratio: 0.44 [95% CI: 0.25, 0.77]; P = 0.004). For EORTC QLQ-C30 symptoms and functioning subscales, only nausea/vomiting symptom score was worse in the olaparib arm than in the TPC arm (across all visits compared with baseline).

Conclusion: HRQoL was consistently improved for patients treated with olaparib, compared with chemotherapy TPC.

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daily) or single-agent, pre-defined, chemotherapy TPC (capecitabine, eribulin or vinorelbine — see Supplementary Materials for dosage).

2.2. Study outcome measures

Primary and secondary efficacy and safety outcomes are described fully in the primary manuscript [5]. The pre-specified PROs reported here were assessed using the paper-based 30-item EORTC QLQ-C30 questionnaire completed by the patient at baseline (before randomisation once eligibility was confirmed) and every 6 weeks until investigator-assessed objective disease progression. Patients who discontinued treatment because of toxicity continued PRO assessments until disease progression. The EORTC QLQ-C30 includes a two-item global QoL scale (global health status/QoL score); five multi-item functional scales (physical, role, emotional, cognitive and social); three multi-item symptom scales (fatigue, pain and nausea/vomiting); five single items assessing common cancer symptoms (dyspnoea, insomnia, appetite loss, constipation and diarrhoea) and a single item addressing the financial impact of disease [8]. Scores for the QLQ-C30 range from 0 to 100. For global health status/QoL score and functional scales, higher scores indicate better HRQoL and level of functioning. For symptom scales, higher scores indicate greater severity of symptoms. The impact of olaparib on symptoms and HRQoL, as assessed by the multi-item functional scales, multi-item symptom scales and common cancer symptoms, were exploratory analyses.

The primary HRQoL end-point compared mean change from baseline between treatment arms in the two-item global health status/QoL score across all visits. Secondly, we assessed the proportion of patients in each arm who experienced a clinically meaningful increase (improvement) or decrease (deterioration) in global health status/QoL score (defined as ≥10-point change from baseline) [9]. Best overall HRQoL response was defined as the best HRQoL response the patient achieved from randomisation to disease progression. Best HRQoL response was categorised as ‘improved’, ‘no change’, ‘deterioration’ or ‘other’ according to the following criteria: ‘improved’ — two visit responses of ‘improved’ sustained for ≥21 days with no intervening response of ‘deterioration’; ‘no change’ — two visit responses of either ‘no change’ or ‘improved’ and ‘no change’ ≥21 days apart with no intervening response of ‘deterioration’; ‘deterioration’ — a visit response of ‘deterioration’ without a response of ‘improved’ or ‘no change’ within 21 days; ‘other’ — patients who met the criteria for ‘improved’, ‘no change’ or ‘deterioration’.

For additional information on continued treatment effectiveness, an analysis of the number of patients remaining on treatment for ≥6 months and time to subsequent therapies are shown in Supplementary Materials.

2.3. Statistical analysis

Sample size determination has been reported in the primary manuscript [5]. Questionnaire compliance and completion data were analysed overall and by study visit and summarised by treatment. All PRO data were analysed using the full analysis set on an intention-to-treat basis that included all randomised patients with a baseline and ≥1 post-baseline assessment. For the primary HRQoL analysis, a linear mixed-model repeated-measures (MMRM) analysis adjusting for score at baseline, time and treatment-by-time interaction was used to estimate the cumulative effect of olaparib versus TPC on global health status/QoL. MMRM methodology uses observed data to implicitly impute unobserved data; by including covariates in the model, the analysis assumes that patients with missing data would behave similarly to other patients with similar values for these covariates (e.g. treatment group) had they not missed the assessment. The analysis included all post-baseline visits up to the last scheduled visit in which ≥20 patients in each treatment arm had an evaluable score. Differences between treatment groups were compared using adjusted mean estimates per treatment group and corresponding 95% CIs and P values.

Best overall global health status/QoL response rates (improvement, no change or deterioration) were summarised descriptively for patients. Multi-item functional and symptom subscales and single-item cancer symptoms were exploratory variables. For global health status/QoL and multi-item functional subscales, best overall response was evaluated in patients with baseline score ≥10. For multi-item symptom subscales and single-item symptom scales, best overall response was evaluated in patients with baseline score ≤90.

Time to deterioration (TTD) of global health status/QoL (planned analysis) and functional and symptom scales (post hoc analysis) was defined as time from randomisation until the date of a clinically important deterioration in the global health status/QoL score or functional and symptom scores. For multi-item functional and global health status/QoL subscales, deterioration included patients with baseline score ≥10; for symptom subscales/single items, deterioration included patients with baseline score ≤90. Clinically important deterioration was defined as a decrease from baseline of ≥10 points (or an increase from baseline of ≥10 points for the symptom scales) that was sustained at the next scheduled visit. Patients who did not experience deterioration by the time of progression (PRO data beyond progression were not collected) were censored at the time of the last-available PRO assessment. Data were analysed using a log-rank test, with HR and 95% CI generated from the log-rank test statistics. The Kaplan–Meier method was used to estimate medians.
3. Results

3.1. Study population characteristics

In total, 302 patients (intention-to-treat population) were randomly assigned to treatment (olaparib, n = 205; TPC, n = 97) between 7 April 2014 and 27 November 2015. Of the 97 TPC patients, 91 received study treatment (capecitabine, n = 41; eribulin, n = 34; vinorelbine, n = 16). A participant flow diagram (Fig. 1) and baseline demographic characteristics, which were well balanced between the two treatment groups, have been described [5]. Mean (standard deviation [SD]) global health status/QoL score (scale range: 0–100; higher scores indicate higher levels of function and QoL) at baseline was 63.2 (21.0) for the olaparib treatment arm and 63.3 (21.2) for TPC (capecitabine 62.3 [19.6], eribulin 59.1 [22.1], vinorelbine 67.7 [21.5]) [5]. Multi-item symptom scores at baseline in both treatment arms were similar to those cited for recurrent or mBC in the EORTC QLQ-C30 reference value manual [10].

3.2. Questionnaire compliance/completion

Compliance and completion rates for EORTC QLQ-C30 at baseline were >95% in both arms (Table 1). Completion rates declined faster in the TPC arm.

3.3. Global health status/QoL

The between-treatment comparison (average over time) based on an MMRM model showed a clinically significant improvement from baseline in mean (SD) global health status/QoL score for olaparib versus TPC (3.9 [1.2] vs –3.6 [2.2]; difference 7.5 [95% CI: 2.48–12.44]; P = 0.0035) [5]. The adjusted mean change from baseline over time (Fig. 2) showed that, for each visit, patients in the olaparib arm had an improvement in mean global health status/QoL score, whereas patients in the TPC arm had a decline.

Median TTD in global health status/QoL (≥10 points) was not reached in the olaparib arm and was 15.3 months in the TPC arm (HR: 0.44; 95% CI: 0.25–0.77; P = 0.004; Fig. 3) [5]. Global scores had deteriorated in fewer patients in the olaparib arm versus TPC at 6 months (18.5% vs 38.8%) and 12 months (36.0% vs 46.5%). Censoring of most olaparib-treated patients because they did not meet the 10-point threshold criterion for deterioration means that these TTD data should be interpreted with caution (see Discussion).

More patients in the olaparib treatment arm showed an improvement in best overall response rates for global health status/QoL versus TPC (33.7% vs 13.4%), and more patients in the TPC arm showed a deterioration in global health status/QoL versus olaparib (20.6% vs 11.7%) (Fig. 4).

3.4. Functional subscales

Kaplan–Meier estimates of TTD (before or at progression) showed that olaparib delayed TTD versus TPC for each of the five functional subscales (Fig. 5). The best overall response data for the functional subscales support the results for global health status/QoL and highlight that, across all five functional

Fig. 1. Participant flow diagram. bid, twice daily; TPC, treatment of physician’s choice.
subscales, more patients experienced an improvement in functional subscales in the olaparib treatment arm versus TPC (Supplementary Fig. 1A); concordantly, more patients experienced deterioration in functional subscales in the TPC arm versus olaparib.

### 3.5. Symptom subscales

For fatigue, pain and nausea/vomiting multi-item symptom scales (decrease in score indicates improvement in symptoms), as well as single-item symptom scales, a higher proportion of patients in the olaparib treatment arm experienced improvements versus TPC (Supplementary Fig. 1B and 1C).

Adjusted mean changes from baseline in symptom scores across all visits (Fig. 6) showed improvements in fatigue, pain, dyspnoea, insomnia, loss of appetite, constipation and diarrhoea scores among patients receiving olaparib versus patients receiving TPC. Only mean nausea/vomiting scores were better (relative to baseline) in the TPC arm versus olaparib. Apart from nausea/vomiting, the HR for TTD favoured olaparib versus TPC across all symptom scales (Supplementary Fig. 2).

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**Table 1**

Compliance and completion rates for the EORTC QLQ-C30 questionnaire.

<table>
<thead>
<tr>
<th>Time point</th>
<th>Expected forms, n</th>
<th>Compliance rate, %</th>
<th>Completion rate, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Olaparib</td>
<td>TPC</td>
<td>Olaparib (n = 205)</td>
</tr>
<tr>
<td>Baseline</td>
<td>205</td>
<td>97</td>
<td>99.0</td>
</tr>
<tr>
<td>Visit 6 (week 6)</td>
<td>205</td>
<td>96</td>
<td>87.3</td>
</tr>
<tr>
<td>Visit 8 (week 12)</td>
<td>184</td>
<td>73</td>
<td>87.0</td>
</tr>
<tr>
<td>Visit 10 (week 18)</td>
<td>167</td>
<td>50</td>
<td>84.4</td>
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<tr>
<td>Visit 12 (week 24)</td>
<td>136</td>
<td>36</td>
<td>84.6</td>
</tr>
<tr>
<td>Visit 14 (week 30)</td>
<td>115</td>
<td>30</td>
<td>84.4</td>
</tr>
<tr>
<td>All visits (overall)</td>
<td>205</td>
<td>97</td>
<td>93.2</td>
</tr>
</tbody>
</table>

**EORTC QLQ-C30**, European Organisation for Research and Treatment of Cancer Quality of Life Questionnaire Core 30-item module; PRO, patient-reported outcome; TPC, treatment of physician’s choice.

Note that 19 assessments are included whereby the questionnaire was administered to a caregiver, relative or unspecified person, rather than directly to the patient.

a Compliance rate for each treatment arm was defined as follows: (the number of evaluable questionnaires [defined as a questionnaire with enough responses to score at least one scale/domain])/(the number of patients expected to complete questionnaires at each visit [i.e. patients still under PRO follow-up]) × 100%.

b Completion rate for each treatment arm was defined as follows: [(the number of received questionnaires)/(PRO analysis population)] × 100%.

c For overall, patients were counted as received/evaluable if they had a received/evaluable baseline and at least one received/evaluable post-baseline form.
3.6. Time to subsequent therapies

Results of the analyses of time to subsequent therapies are presented in Supplementary Materials.

4. Discussion

Optimising HRQoL is a key component in the guidelines for treatment and management of mBC as
treatment is usually not curative but palliative [1]. Our findings show that the efficacy benefits reported with olaparib among patients with gBRCAm HER2-negative mBC are accompanied by improvements in symptoms, functioning and HRQoL assessed using the EORTC QLQ-C30.

Adjusted mean change from baseline in global health status/QoL score across all visits revealed greater improvements with olaparib versus TPC. Overall, HRQoL in patients receiving olaparib improved during time on treatment versus patients receiving TPC, in whom deterioration in HRQoL over time was observed. The difference between arms is consistent with a small, clinically significant improvement (7.5-point difference [moderate, 10–20; large, >20]) on a group-level assessment [9,11]. TTD was also extended in the olaparib arm versus TPC. However, the median TTD findings should be interpreted cautiously as PRO data were collected only until disease progression, and most patients had not reached the 10-point threshold criterion for deterioration at this time. Patients who did not meet the deterioration criteria were censored at the time of their last global health status/QoL assessment. Given these limitations, the best overall response represents a more instructive assessment to determine the impact of treatments on HRQoL. The findings of the best overall response showed that olaparib-treated patients were almost three times as likely as those receiving TPC to report a clinically significant improvement in global health status/QoL.

EORTC QLQ-C30 functional and symptom subscale scores were generally improved in the olaparib arm versus TPC. Marked improvements in functioning favoured olaparib over TPC, as measured on the physical, cognitive and emotional subscales. For each symptom subscale, more patients in the olaparib group showed a best overall response of a clinically meaningful improvement in symptom score versus TPC. Single symptoms included in the scale are considered as being important to patients with breast cancer. Our findings showed particular benefits of olaparib over TPC in terms of improved pain and dyspnoea. For patients who experienced deterioration in symptoms, this was typically greater for those receiving TPC versus olaparib.

Differences in adjusted mean change from baseline in symptom scores for nausea/vomiting favoured TPC. This is a known effect of olaparib that has been shown to be greater during treatment initiation [12]. Indeed, a high proportion of patients receiving olaparib reported an improvement in the nausea/vomiting subscale over time. Previously reported safety data from the OlympiAD trial showed that more patients experienced nausea/vomiting with olaparib versus TPC, but no grade ≥3 events were reported and discontinuation rates were low [5,13]. We found that although a higher proportion of olaparib-treated patients reported a best overall response of improvement in nausea/vomiting, differences in adjusted mean change from baseline in symptom scores for nausea/vomiting favoured TPC. These data are hard to reconcile but suggest that patients whose symptoms do not improve experience a greater decrease in score. In clinical practice, supportive measures such as early antiemetic administration may be appropriate to manage events of nausea and vomiting, thus minimising olaparib dose adjustments/discontinuations. The improvements in HRQoL in the olaparib arm versus TPC are consistent with the lower rate of discontinuations because of adverse events (4.9% vs 7.7%) and the longer median treatment duration (8.2 months) reported previously in the OlympiAD trial [5]. The supportive analyses of time to subsequent therapies (see Supplementary Materials) further expand on these data, showing that more patients in the olaparib arm remained on treatment for ≥6 months versus TPC (60.0% vs 27.5%), and median time to first and second subsequent therapy was longer with olaparib versus TPC (9.4 vs 4.2 months and 14.3 vs 10.5 months, respectively); time on assigned treatment has been recognised as a meaningful clinical trial end-point in settings whereby a disease course involves multiple rounds of subsequent treatment after first progression [14].

Limitations of the study include the open-label trial design, which has the potential to introduce patient biases. In addition, the absence of post-progression PRO data collection may have underestimated the impact of olaparib on HRQoL given that patients receiving TPC had disease progression earlier than those receiving olaparib, and progression would be expected to result in a further decrease in HRQoL. However, this would also be dependent on the HRQoL impact of post-progression treatment. Questionnaire compliance and completion rates were lower in the TPC arm versus olaparib (76.3% vs 93.2%), and completion rates also declined faster in the TPC arm because of earlier progression and cessation of EORTC QLQ-C30 data collection. This higher rate of missing data may also have biased the results of the analysis. The direction of any bias on the effect of treatment is unclear as it depends on whether missing data would have improved or worsened PRO measures for the TPC arm. However, the selective aspect of completing EORTC QLQ-C30 data collection is likely to be largely explained by observed covariates included in the MMRM analysis, i.e. treatment arm and baseline HRQoL score, which assumes that patients with missing data would behave similarly to other patients in the same treatment group, and with similar covariate values, had they not missed the
Fig. 5. Kaplan–Meier estimates of time to deterioration for EORTC QLQ-C30 functional subscales before or at progression. (a) Physical; (b) role; (c) social; (d) cognitive and (e) emotional. bid, twice daily; CI, confidence interval; EORTC QLQ-C30, European Organisation for Research and Treatment of Cancer Quality of Life Questionnaire Core 30-item module; HR, hazard ratio; NC, not calculable; TPC, treatment of physician’s choice.
assessment. Studies have shown that when such observed covariates are included, the adjusted estimates are robust even when data are not missing completely at random [15].

Favourable PROs have also been reported with another PARP inhibitor among patients with gBRCAm advanced breast cancer [6]. Improvement in the global health score from baseline with talazoparib in the EMBRACA trial (3.0) was similar to that reported here (3.9). Our findings and those of EMBRACA provide further support of the beneficial effects of PARP inhibitors on HRQoL versus chemotherapy in patients with mBC. Based on these data, recently updated advanced breast cancer (ABC4) and National Comprehensive Cancer Network guidelines recommend PARP inhibitors as a treatment option for BRCA-associated
5. Conclusions

The planned and post hoc analyses of data we report from the phase III OlympiAD study suggest that olaparib treatment can lead to improvements in the symptoms, functioning and HRQoL of patients with gBRCAm HER2-negative mBC versus chemotherapy TPC. PRO results will be valuable to both physicians and patients when considering the clinical benefits of PARP inhibitor treatment in mBC, versus chemotherapy.

Data access, responsibility and analysis

Data underlying the findings described in this manuscript may be obtained in accordance with AstraZeneca’s data sharing policy described at https://astrazenecagrouptrials.pharmacm.com/ST/Submission/Disclosure.

M.R., W.B. and C.G. had full access to all the data in the study and take responsibility for the integrity of the data and the accuracy of the data analysis. W.B. is the statistical author who has responsibility for the statistical analysis.

Conflict of interest statement

M.R. reports receiving honoraria (advisory) from AstraZeneca and has served a consulting or advisory role for AstraZeneca, Daiichi Sankyo (uncompensated), McKesson, Merck (uncompensated) and Pfizer (uncompensated). Research funding has been received from AbbVie (institution), AstraZeneca (institution), Invitae (institution, in-kind), Medivation (institution), Myriad Genetics (institution, in-kind), Pfizer (institution) and Tesaro (institution). Travel or accommodation expenses have been received from AstraZeneca, and other transfer of value items are from AstraZeneca (editorial services) and Pfizer (editorial services). K.J.R. reports intellectual property interests for herself and her spouse regarding a discovery or technology relating to health or medicine. S.-A.I. has served a consultant role for AstraZeneca, Novartis, Spectrum, Hanmi, Pfizer and Roche and reports receiving research funding from AstraZeneca. E.S. reports receiving honoraria from Amgen, Astellas, AstraZeneca, Bayer, Bristol-Myers Squibb, Celgene, Clinigen, Egis, Eli Lilly, Janssen, Novartis, Pfizer, Pierre Fabre, priME, Roche and Teva; travel support from Amgen, AstraZeneca, Egis, Novartis, Pfizer and Roche and research funding from Amgen, Astellas, AstraZeneca, Bayer, Bristol-Myers Squibb, Boehringer, Eli Lilly, Janssen, Merck, Novartis, Pfizer, Roche and Samsung. B.X. reports no conflicts of interest. S.M.D. reports receiving honoraria from AstraZeneca, Clovis Oncology and Bristol-Myers Squibb, and research funding to the University of Pennsylvania from AstraZeneca and Clovis Oncology. N.M. reports receiving honoraria from Chugai Pharma, AstraZeneca, Pfizer and Takeda, and institutional research funding from Chugai Pharma, AstraZeneca, Kyowa Hakka Kirin, MSD, Novartis, Pfizer, Eli-Lilly and Daiichi Sankyo. W.L. reports no conflicts of interest.
interest. N.T. reports consulting for AstraZeneca and receiving grant support from Myriad Genetics and Ambry Genetics. A.A. reports receiving fees for serving on an advisory board from Roche and Syndax Pharmaceuticals and her spouse holding stock options in AstraZeneca. S.D. reports receiving honoraria from Novartis, Roche, AstraZeneca, Pfizer, GE Healthcare and Puma Biotechnology; consulting fees for Novartis, Roche, Pfizer, AstraZeneca and Puma Biotechnology; research funding from Novartis, Roche, AstraZeneca, Pfizer and Puma Biotechnology; and travel and accommodation expenses from Pfizer, AstraZeneca and Novartis. W.B., C.G., A.D. and R.H. are employees of AstraZeneca. P.C. reports being a member of speakers’ bureaus of Roche, Novartis, AstraZeneca and GlaxoSmithKline and receiving travel and accommodation expenses from Novartis, GlaxoSmithKline and Celgene, and research funding from Roche, Novartis and Merck Serono.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ejca.2019.06.023.

References