



#### **ORIGINAL ARTICLE**

## Overall survival in the OlympiA phase III trial of adjuvant olaparib in patients with germline pathogenic variants in BRCA1/2 and high-risk, early breast cancer

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Background: The randomized, double-blind OlympiA trial compared 1 year of the oral poly(adenosine diphosphate-ribose) polymerase inhibitor, olaparib, to matching placebo as adjuvant therapy for patients with pathogenic or likely pathogenic variants in germline *BRCA1* or *BRCA2* (*gBRCA1/2pv*) and high-risk, human epidermal growth factor receptor 2-negative, early breast cancer (EBC). The first pre-specified interim analysis (IA) previously demonstrated statistically significant improvement in invasive disease-free survival (IDFS) and distant disease-free survival (DDFS). The olaparib group had fewer deaths than the placebo group, but the difference did not reach statistical significance for overall survival (OS). We now report the pre-specified second IA of OS with updates of IDFS, DDFS, and safety.

**Patients and methods:** One thousand eight hundred and thirty-six patients were randomly assigned to olaparib or placebo following (neo)adjuvant chemotherapy, surgery, and radiation therapy if indicated. Endocrine therapy was given concurrently with study medication for hormone receptor-positive cancers. Statistical significance for OS at this IA required P < 0.015.

**Results:** With a median follow-up of 3.5 years, the second IA of OS demonstrated significant improvement in the olaparib group relative to the placebo group [hazard ratio 0.68; 98.5% confidence interval (CI) 0.47-0.97; P=0.009]. Four-year OS was 89.8% in the olaparib group and 86.4% in the placebo group ( $\Delta$  3.4%, 95% CI -0.1% to 6.8%). Four-year IDFS for the olaparib group versus placebo group was 82.7% versus 75.4% ( $\Delta$  7.3%, 95% CI 3.0% to 11.5%) and 4-year DDFS was 86.5% versus 79.1% ( $\Delta$  7.4%, 95% CI 3.6% to 11.3%), respectively. Subset analyses for OS, IDFS, and DDFS demonstrated benefit across major subgroups. No new safety signals were identified including no new cases of acute myeloid leukemia or myelodysplastic syndrome.

**Conclusion:** With 3.5 years of median follow-up, OlympiA demonstrates statistically significant improvement in OS with adjuvant olaparib compared with placebo for gBRCA1/2pv-associated EBC and maintained improvements in the previously reported, statistically significant endpoints of IDFS and DDFS with no new safety signals.

Key words: breast cancer, BRCA1/2, PARP inhibition, olaparib, adjuvant therapy

#### **INTRODUCTION**

Cancers harboring germline pathogenic or likely pathogenic variants in *BRCA1* or *BRCA2* (*gBRCA1/2*pv) are characterized by homologous recombination DNA repair deficiency following the inactivation of the wildtype allele during tumor evolution. This engenders selective sensitivity to inhibition and trapping of the DNA repair enzyme, poly (adenosine diphosphate-ribose) polymerase 1 (PARP1) by exploiting the concept of synthetic lethality, as functional homologous recombination is required for cell survival when PARP1 function is inhibited and PARP1 is trapped on DNA arresting the DNA replication apparatus. Olaparib and talazoparib both inhibit and trap PARP1 on DNA and have been approved for treating patients with *gBRCA1/2*pv and metastatic breast cancer (MBC) irrespective of hormone receptor status. 5,6

Breast cancers associated with gBRCA1/2pv are characterized by high-grade disease with most gBRCA1pv-

associated tumors being triple negative, whereas most gBRCA2pv-associated cancers are hormone receptor positive and human epidermal growth factor receptor 2 (HER2) negative, 7-9 and often associated with high-risk classification on RNA-based prognostic assays. 10,11 Because patients with gBRCA1/2pv-associated early breast cancers (EBCs) and highrisk clinico-pathological features remain at increased risk for recurrence following standard multimodality therapies, OlympiA (ClinicalTrials.gov: NCT02032823) was designed to determine whether 1 year of adjuvant olaparib could improve outcomes in this population. This phase III, doubleblinded, placebo-controlled study randomized 1836 eligible patients with gBRCApv-associated EBC from 2014 to 2019. Following review of the first pre-specified interim analysis (IA1) of the primary endpoint of invasive disease-free survival (IDFS), the independent data monitoring committee (IDMC) recommended full analysis, which was previously reported. 12 With a median follow-up of 2.5 years, patients

randomized to olaparib had statistically significant and clinically meaningful improvement in IDFS compared to placebo [hazard ratio (HR) 0.58; 99.5% CI 0.41-0.82; P < 0.001] and distant disease-free survival (DDFS) (HR 0.57; 99.5% CI 0.39-0.83; P < 0.001), which corresponded to absolute improvements at 3 years in IDFS of 8.8% and in DDFS of 7.1%. The number of deaths in the olaparib group was fewer than in the placebo group (59 versus 86), but the difference (HR 0.68; 99% CI 0.44-1.05; P = 0.02) did not meet the prespecified boundary for statistical significance for overall survival (OS) (P < 0.01). The safety analysis was consistent with the experience in the MBC setting and provided no early evidence of increased risk of acute myeloid leukemia or myelodysplastic syndrome (AML/MDS).  $^{12}$ 

The second IA (IA2) of OS was pre-specified to occur when 330 IDFS events had been reported in the study population. Here we report the results of this OS analysis with updates of IDFS, DDFS, and safety information.

#### **PATIENTS AND METHODS**

#### Study design and patient population

Details of study design and populations for the primary and secondary efficacy endpoints and safety are described in the original manuscript. The trial was conducted in accordance with the amended Declaration of Helsinki and the protocol was approved by the institutional review board at each participating center. All patients provided written informed consent. Olaparib and placebo were provided by AstraZeneca.

In summary, eligible, consenting patients with gBRCA1/2pv determined by germline testing at the site or centrally, with high-risk, HER2-negative EBC were randomized to receive 1 year of study medication consisting of either oral olaparib 300 mg b.i.d. or matching placebo, stratified by hormone receptor status, prior neoadjuvant (NACT) versus adjuvant (ACT) chemotherapy, and platinum therapy for current breast cancer (yes versus no). Eligible patients had received at least six cycles of NACT or ACT containing a taxane, an anthracycline, or both, had completed surgery, and had completed adjuvant radiotherapy if indicated according to local standards at least 2 weeks before randomization. Patients with hormone receptor-positive cancers were to receive at least 5 years of adjuvant endocrine therapy (ET) as per local standards concurrent with study medication. Bisphosphonates and denosumab were allowed as per investigator's discretion. Patients who had received NACT could not receive postoperative chemotherapy.

Eligible patients with triple-negative breast cancer (TNBC) included those who received NACT with residual invasive cancer in the breast or axillary nodes, and those who received ACT were either node positive or node negative with a T2-T4 primary tumor at initial surgery. Following an early amendment, patients with hormone receptor-positive, HER2-negative disease became eligible with a clinical and pathological stage plus estrogen receptor and nuclear grade (CPS + EG) score of  $\geq$ 3 following NACT<sup>14,15</sup> or  $\geq$ 4 positive nodes at initial surgery.

#### **Endpoints and assessments**

In accordance with the Standardized Definitions for Efficacy End Points (STEEP) system,<sup>16</sup> the primary endpoint of IDFS was defined as the time from randomization until the date of first occurrence of one of the following events: ipsilateral invasive breast tumor, locoregional invasive disease, distant recurrence, contralateral invasive breast cancer, second primary invasive cancer, or death from any cause. Patients without a documented IDFS event were censored at the date they were last known to be disease free. Secondary endpoints include DDFS, defined as time from randomization until documented evidence of first distant recurrence of breast cancer or death, and OS defined as time from the date of randomization until death due to any cause.

Efficacy analyses were based on the intention-to-treat (ITT) population. Survival functions were estimated by Kaplan—Meier method. The stratified Cox proportional hazards model was used to estimate the HR and confidence intervals (CIs), and the *P* value for the comparison of survival between treatment arms was generated by stratified log-rank test. Safety was assessed in the population who received at least one dose of study medication.

OlympiA was designed to achieve a 90% power to detect an HR of 0.70 for the primary endpoint of IDFS, assuming a twosided 5% significance level. With a sample size of 1800 patients, the primary analysis of IDFS would be triggered by 330 IDFS events in the ITT population. Four analysis time-points were pre-planned, with a hierarchical multiple testing procedure to strongly control type 1 error across analysis timepoints and endpoints (Supplementary Table S1, available at https://doi.org/10.1016/j.annonc.2022.09.159). As previously reported,<sup>12</sup> the IA of IDFS in the entire ITT population was triggered when 165 IDFS events had been observed in the first 900 patients randomized (IA1). Superiority boundaries were P < 0.005 for IDFS, followed by P < 0.005 for DDFS, and P < 0.01 for OS (Supplementary Table S1, available at https://doi.org/ 10.1016/j.annonc.2022.09.159). Superiority boundaries for both IDFS and DDFS were crossed, but not for OS. 12 The second pre-specified IA2 of OS was triggered by 330 IDFS events in the ITT population and results are presented herein. The boundary for the two-sided significance test of OS at IA2 was P < 0.015; thus, 98.5% CIs for OS are calculated in this analysis. Updated analyses of IDFS and DDFS were carried out with 95% CIs as these endpoint analyses are now descriptive.

#### **RESULTS**

#### **Patients**

From June 2014 through May 2019, 1836 patients were randomly assigned to receive either olaparib or placebo. IA2 was triggered on 12 July 2021; case report forms for study visits up to data cut-off for IA2 were collected and data quality controlled with database lock occurring on 17 December 2021. Median follow-up was 3.5 years [interquartile range (IQR) 2.5-4.5 years] in the ITT population, 3.6 years (IQR 2.5-4.7 years) in the TNBC cohort, and 3.4 years (IQR 2.5-4.1 years) in the hormone receptor-positive cohort.

| Characteristic  | Olaparib ( $n = 921$ ) | Placebo ( $n = 915$ ) |
|---|------------------------|-----------------------|
| Age, median (interquartile range), years              | 42 (36-49)             | 43 (36-50)            |
| gBRCA P/LP gene—n (%) <sup>b</sup>                    |                        |                       |
| BRCA1   | 656 (71.2)             | 669 (73.1)            |
| BRCA2   | 260 (28.2)             | 238 (26.0)            |
| BRCA1 and BRCA2                                       | 2 (0.2)                | 5 (0.5)               |
| No gBRCA P/LP variant                                 | 2 (0.2)                | 3 (0.3)               |
| Missing   | 1 (0.1)                | 0 (0.0)               |
| Prior adjuvant/neoadjuvant chemotherapy, n (%)        |                        |                       |
| Adjuvant  | 461 (50.1)             | 455 (49.7)            |
| Neoadjuvant   | 460 (49.9)             | 460 (50.3)            |
| Anthracycline and taxane regimen                      | 871 (94.6)             | 849 (92.8)            |
| Anthracycline regimen (without taxane)                | 7 (0.8)                | 13 (1.4)              |
| Taxane regimen (without anthracycline)                | 43 (4.7)               | 52 (5.7)              |
| Regimen not reported                                  | 0 (0.0)                | 1 (0.1)               |
| <6 cycles of (neo)adjuvant chemotherapy               | 7 (0.8)                | 13 (1.4)              |
| Platinum-based (neo)adjuvant therapy                  |                        |                       |
| No  | 674 (73.2)             | 677 (74.0)            |
| Yes   | 247 (26.8)             | 238 (26.0)            |
| Concurrent hormone therapy (hormone receptor positive | 146/168 (86.9)         | 146/157 (93.0)        |
| only), n (%)  |                        |                       |
| Hormone receptor status, n (%) <sup>c</sup>           |                        |                       |
| Hormone receptor positive/HER2 negative <sup>d</sup>  | 168 (18.2)             | 157 (17.2)            |
| Triple-negative breast cancer <sup>e</sup>            | 751 (81.5)             | 758 (82.8)            |
| Menopausal status (females only), n (%)               |                        |                       |
| Premenopausal   | 572/919 (62.2)         | 553/911 (60.7)        |
| Postmenopausal  | 347/919 (37.8)         | 358/911 (39.3)        |
| Primary breast cancer surgery, n (%)                  |                        |                       |
| Mastectomy  | 699 (75.9)             | 674 (73.7)            |
| Conservative surgery only                             | 222 (24.1)             | 239 (26.1)            |
| Missing   | 0 (0.0)                | 2 (0.2)               |

HER2, human epidermal growth factor receptor 2; P/LP, pathogenic or likely pathogenic variants

<sup>a</sup>Further information on baseline characteristics is provided in Supplementary Table S2, available at https://doi.org/10.1016/j.annonc.2022.09.159. Percentages may not total 100 because of rounding.

<sup>b</sup>For a detailed description of local and central Myriad BRCA testing in patients enrolled in the trial, see Supplementary Figure S2, available at https://doi.org/10.1016/j.annonc. 2022.09.159. Variant interpretation by Myriad Genetics (BRCAnalysis) (1649 patients) and BGI Genomics (247 patients) was carried out with the use of multiple established databases (e.g. ClinVar, ClinGen, and ENIGMA) and published and internal functional and clinical data, compliant with American College of Medical Genetics published guidelines. Eighty-five patients randomized in China had variant interpretation by both BGI Genomics and Myriad Genetics. The 24 pathogenic or likely pathogenic variants from local laboratories without central Myriad confirmation were confirmed by the OlympiA genetics advisory committee with the use of published databases as above. Discordant data are referred to in Supplementary Figure S2, available at https://doi.org/10.1016/j.annonc.2022.09.159. Listing of pathogenic or likely pathogenic *BRCA1* and *BRCA2* variants that occurred in more than one patient have previously been reported.<sup>12</sup>

<sup>c</sup>Hormone receptor status was defined by local test results.

<sup>d</sup>The original protocol that was activated in 2014 was developed for HER2-negative patients but included only patients with triple-negative breast cancer after regulatory review. When the safety rationale with respect to recurrence risk relative to combination therapy with olaparib and endocrine therapy was accepted by regulators, the protocol was amended in 2015 to include patients with high-risk hormone receptor-positive disease and to increase the sample size to the current number of 1800 patients (see the protocol). The first patient with hormone receptor-positive disease was enrolled in December 2015.

 $^{
m e}$ Triple-negative breast cancer was defined in the eligibility criteria as estrogen receptor negative and progesterone receptor negative, as indicated by immunohistochemical (IHC) nuclear staining of <1%, and HER2 negative (not eligible for anti-HER2 therapy), as indicated by one of the following: an IHC score of 0 or 1+; an IHC score of 2+ and HER2-nonamplified disease on in situ hybridization (ISH) with a ratio of <2.0 and, if reported, an average HER2 copy number of <4 signals per cell; or HER2-nonamplified disease on ISH with a ratio of <2.0 and, if reported, an average HER2 copy number of <4 signals per cell (without IHC). Two patients (both in the olaparib group) were excluded from the summary of the subgroup with triple-negative breast cancer because they did not have confirmed HER2-negative status.

After randomization, 10 patients in the olaparib group and 11 in the placebo group did not receive assigned therapy (Supplementary Figure S1: Consort Diagram, available at https://doi.org/10.1016/j.annonc.2022.09.159). Baseline characteristics of the patients were balanced between the two treatment groups (Table 1, Supplementary Table S2, available at https://doi.org/10.1016/j.annonc.2022.09.159). Most of the patients (82.2%) had TNBC. Approximately half of them received ACT and half NACT, with the majority (93.7%) receiving both an anthracycline and a taxane. A platinum agent was also received by 26.4% of patients, primarily in the NACT setting. Germline *BRCA1*pv were present in 72.2% and *gBRCA2*pv in 27.1% of patients with

even distribution between treatment groups. Seven patients had both gBRCA1pv and gBRCA2pv.

#### **Efficacy**

OS was significantly improved in the olaparib group relative to the placebo group (HR 0.68; 98.5% CI 0.47-0.97; P=0.009) (Figure 1A). Deaths were now reported in 75 patients (8.1%) in the olaparib group and 109 (11.9%) in the placebo group, 16 and 23 more, respectively, than at the previous IA. The cause of death was breast cancer in 93.3% in the olaparib group and 94.5% in the placebo group (Supplementary Table S3, available at https://doi.org/10.1016/j.annonc.2022.09.159). Death without a prior IDFS

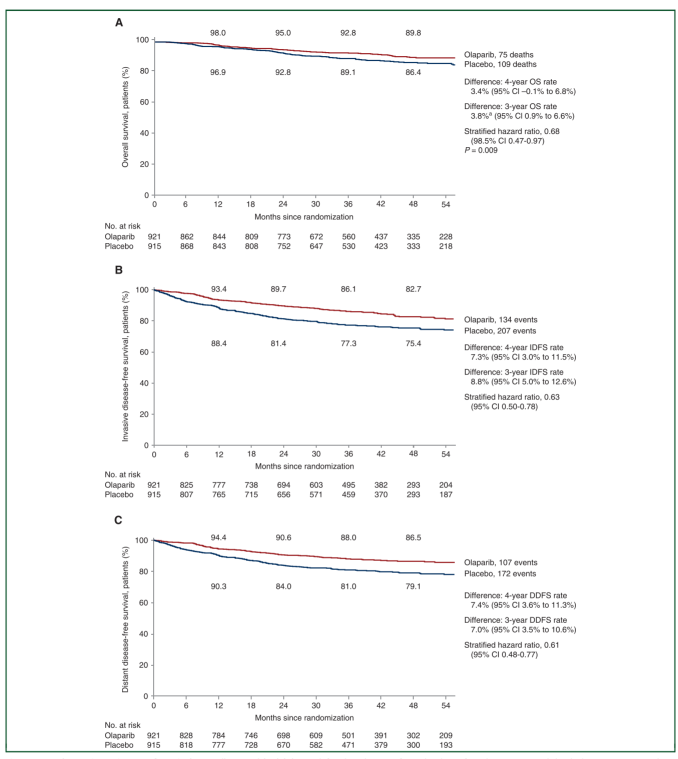


Figure 1. Kaplan-Meier Estimates of Survival. Overall survival (OS) (A) was defined as the time from the date of randomization until death due to any cause; the *P* value for the boundary for significance in this prespecified event-driven interim analysis was <0.015. In accordance with the standardized definitions for efficacy end points (STEEP) system, the primary end point of invasive disease-free survival (IDFS) (B) was defined as the time from randomization until the date of one of the following events: ipsilateral invasive breast tumor, locoregional invasive disease, distant recurrence, contralateral invasive breast cancer, second primary invasive cancer, or death from any cause. Data for patients without a documented event of invasive disease or death were censored at the date they were last known to be disease-free. Distant disease-free survival (DDFS) (C) was defined as the time from randomization until documented evidence of first distant recurrence of breast cancer or death. Distant recurrence includes the following events: distant recurrence (metastatic breast cancer that has either been biopsy confirmed or radiologically diagnosed as recurrent invasive breast cancer); death attributable to any cause, including breast cancer, non-breast cancer, or unknown cause; and second primary non-breast invasive cancer. Evidence of distant recurrence requires either radiologic examination or histopathological confirmation by biopsy. For IDFS and DDFS, 95% confidence intervals only are shown for the hazard ratios, as these results are descriptive. Similarly, the 98.5% confidence interval is shown for the hazard ratio for OS because a *P* value of <0.015 is required to indicate statistical significance for OS. On the basis of the pooling strategy for stratification factors described in Section 2 in the Supplementary Appendix, the primary stratified Cox proportional hazards model of IDFS, DDFS, OS, and the stratified log-rank test of OS, were based on the stratification factor of hormone receptor status only. The e

event was reported in two patients in the olaparib group: one with cardiac arrest and one of unknown cause (Supplementary Table S4, available at https://doi.org/10. 1016/j.annonc.2022.09.159). The percentage of patients alive at 4 years from randomization was 89.8% in the olaparib group and 86.4% in the placebo group (3.4% difference: 95% CI -0.1% to 6.8%) (Figure 1A).

Planned subgroup analyses of OS demonstrated point estimates for improved OS for olaparib consistent with that of the overall population across stratification and gBRCA1pv or gBRCA2pv groups (Figure 2A). The survival benefit of olaparib was observed irrespective of gBRCA1pv or gBRCA2pv groups, hormone receptor status, prior platinum use, and ACT versus NACT context, with Cls that include the point estimate of the HR for OS in the overall population. There was no evidence of statistical heterogeneity in the treatment effect for OS across the subgroups analyzed. Consistent results were also noted in three pre-specified sensitivity analyses of OS described in the Supplementary Methods, available at https://doi.org/10.1016/j.annonc.2022.09.159, and shown in Supplementary Table S5, available at https://doi.org/10.1016/j.annonc.2022.09.159.

With ∼1 year of additional median follow-up, the improvement in the primary endpoint of IDFS observed at the initial analysis 12 was sustained with a similar treatment effect size observed: HR 0.63; 95% CI 0.50-0.78 (Figure 1B). The event frequency of all categories of IDFS events remained lower with olaparib. Distant recurrence comprised 88/134 (65.7%) of IDFS events in the olaparib group and 136/207 (65.7%) in the placebo group (Supplementary Table S4, available at https://doi.org/10. 1016/j.annonc.2022.09.159). IDFS at 4 years was 82.7% in the olaparib group and 75.4% in the placebo group (7.3%) difference: 95% CI 3.0% to 11.5%) (Figure 1B). DDFS was improved in patients who received olaparib (HR 0.61; 95% CI 0.48-0.77). DDFS at 4 years was 86.5% in the olaparib group and 79.1% in the placebo group (7.4% difference: 95% CI 3.6% to 11.3%) (Figure 1C).

Subgroup analysis of IDFS across stratification and gBRCA1pv or gBRCA2pv groups revealed point estimates of treatment effect favoring olaparib over placebo consistent with that of the overall analysis population (Figure 2B). The benefit of adjuvant olaparib relative to placebo was observed irrespective of gBRCA1pv or gBRCA2pv groups, hormone receptor status, prior platinum use, and ACT versus NACT context, with Cls that include the point estimate of the HR for IDFS in the overall population. Update of previously reported detailed subgroup analyses of IDFS<sup>12</sup> is provided in Supplementary Table S6, available at https://doi.org/10.1016/j.annonc.2022.09.159. Subgroup analyses of DDFS across stratification and gBRCA1pv or gBRCA2pv groups revealed similar findings (Figure 2C).

#### Safety

At this safety analysis all patients had completed the protocol-specified course of olaparib or placebo which included 1815 patients (911 in the olaparib group and 904

in the placebo group). The median exposure duration was 364 days on olaparib and 365 days on placebo (Supplementary Table S7, available at https://doi.org/10. 1016/j.annonc.2022.09.159), with median percentage of intended dose delivered being 94.5% in the olaparib group and 98.9% in the placebo group (Supplementary Table S8, available at https://doi.org/10.1016/j.annonc.2022.09.159). Greater than 11 months of the planned 12 months of therapy was completed by 76.1% of patients receiving olaparib compared to 81.7% on placebo (Supplementary Table S9, available at https://doi.org/10.1016/j.annonc. 2022.09.159). In the olaparib group, 228 patients (25.0%) required a dose reduction compared to 47 (5.2%) in the placebo group (Supplementary Table S10, available at https://doi.org/10.1016/j.annonc.2022.09.159). Dose interruptions lasting at least 3 days occurred in 405 (44.5%) patients in the olaparib group and 279 (30.9%) in the placebo group (Supplementary Table S11, available at https:// doi.org/10.1016/j.annonc.2022.09.159). Adverse events (AEs) requiring permanent discontinuation of the trial drug occurred in 98 patients (10.8%) in the olaparib group and 42 (4.6%) in the placebo group. The most frequent AEs leading to discontinuation of olaparib were nausea (2.2%), anemia (1.8%), fatigue (1.6%), and neutrophil count decreased (1%) (Supplementary Table S12, available at https://doi.org/10. 1016/j.annonc.2022.09.159).

Key AE categories are updated and summarized in Table 2 and Supplementary Table S13, available at https://doi.org/ 10.1016/j.annonc.2022.09.159. AEs of any grade with an incidence of ≥10% are updated in Supplementary Table S14, available at https://doi.org/10.1016/j.annonc. 2022.09.159. Grade 3 or higher AEs occurring in >1% of patients were anemia (8.7%), neutropenia (4.9%), leukopenia (3.0%), fatigue (1.8%), and lymphopenia (1.3%), all in the olaparib group. Serious AEs (SAEs) occurred in 79 patients (8.7%) who received olaparib, and 78 (8.6%) who received placebo. AEs leading to death were cardiac arrest in one patient receiving olaparib, and acute myeloid leukemia (AML) and ovarian cancer each in one patient receiving placebo (Table 2). Red blood cell (RBC) transfusion requirements were previously reported<sup>12</sup> and final updates are provided in Supplementary Table S15A and B, available at https://doi.org/10.1016/j.annonc.2022.09.159.

AEs of special interest (AESI) included pneumonitis, radiation pneumonitis, AML/MDS, and new primary malignancies other than AML/MDS. None of the categories had more AESI reported with olaparib relative to placebo (Supplementary Table S13, available at https://doi.org/10.1016/j.annonc.2022.09.159). As of the primary analysis, there were two cases of MDS/AML reported in the olaparib group and three in the placebo group. With additional follow-up, no additional cases of AML or MDS have been reported in either arm.

#### DISCUSSION

The pre-specified second IA of OS in the OlympiA trial demonstrates that 1 year of adjuvant olaparib relative to placebo

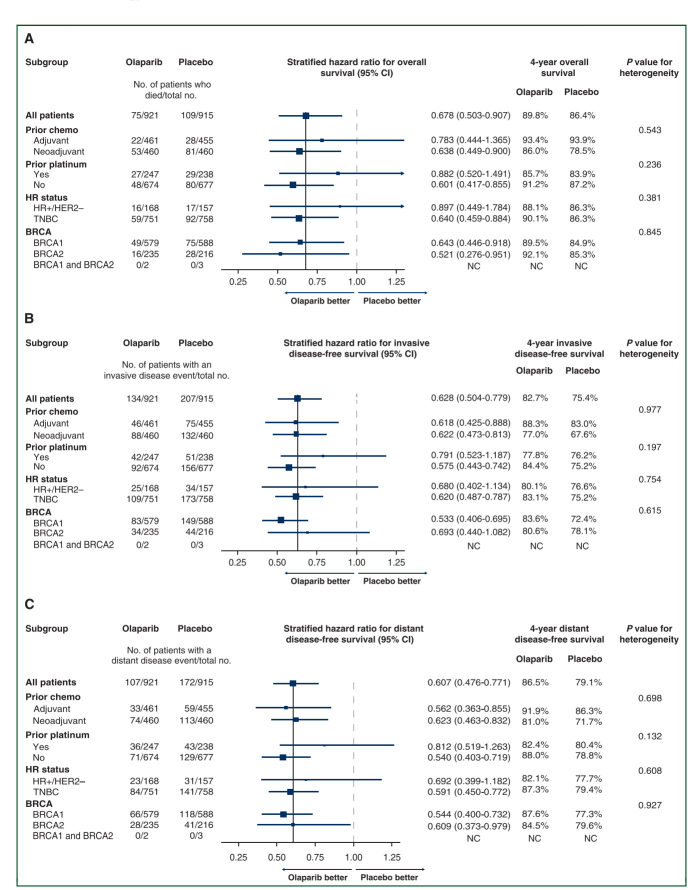


Figure 2. Subgroup analyses by stratification factors and gBRCA1pv or gBRCA2pv groups. (A-C) The solid vertical line indicates the overall hazard-ratio estimate, and the dashed vertical line indicates a hazard ratio of 1.00, as recommended by Cuzick (Cuzick J. Forest plots and the interpretation of subgroups. Lancet 2005; 365:1308). The size of the blue squares corresponds to the number of events contributing to the estimate of the treatment effect. Even without correcting for multiple comparisons, none of the tests for heterogeneity reached statistical significance. BRCA mutation data reflect central Myriad testing results only.

| Table 2. Summary of adverse events in the safety analysis set <sup>a</sup> |            |            |  |
|--|------------|------------|--|
| Adverse event, no. of patients (%)  Olaparib Placebo (n = 911) (n = 904)   |            |            |  |
| Any adverse event  | 836 (91.8) | 758 (83.8) |  |
| Serious adverse event  | 79 (8.7)   | 78 (8.6)   |  |
| Adverse event of special interest <sup>b</sup>                             | 31 (3.4)   | 51 (5.6)   |  |
| MDS/AML  | 2 (0.2)    | 3 (0.3)    |  |
| Pneumonitis <sup>c</sup>   | 9 (1.0)    | 12 (1.3)   |  |
| New primary malignancy <sup>d</sup>  | 21 (2.3)   | 36 (4.0)   |  |
| Grade ≥3 adverse event   | 223 (24.5) | 102 (11.3) |  |
| Grade 4 adverse event <sup>e</sup>   | 17 (1.9)   | 4 (0.4)    |  |
| Adverse event leading to permanent discontinuation of treatment f          | 98 (10.8)  | 42 (4.6)   |  |
| Adverse event leading to death <sup>g</sup>                                | 1 (0.1)    | 2 (0.2)    |  |

AML, acute myeloid leukemia; MDS, myelodysplastic syndrome

<sup>a</sup>Includes adverse events with an onset date on or after the first dose date and up to and including 30 days following date of the last dose of study medication.

<sup>b</sup>Includes adverse events of special interest with onset at any date after first dose of olaparib or placebo. One patient in the olaparib group had both pneumonitis and a new primary invasive breast cancer and is counted in both the pneumonitis and new primary cancer categories.

<sup>c</sup>In the olaparib group, seven patients had pneumonitis, and two patients had radiation pneumonitis. In the placebo group, eight patients had pneumonitis, and four patients had radiation pneumonitis.

<sup>d</sup>Detailed information on the number of patients in each group with specific new primary cancers is provided in Supplementary Table S13, available at https://doi.org/10.1016/j.annonc.2022.09.159.

<sup>e</sup>A total of 18 grade 4 adverse events were reported in 17 patients who received olaparib; 1 patient had both grade 4 anemia and decreased neutrophil count. In the olaparib group, grade 4 adverse events included decreased neutrophil count (in five patients), anemia (in four patients), decreased lymphocyte count (in three patients), and AML, bipolar disorder, fatigue, febrile neutropenia, abnormal hepatic function, and a suicide attempt (in one patient each). In the placebo group, grade 4 adverse events included depression (in two patients) and increased aspartate aminotransferase level and acute cholecystitis (in one patient each).

<sup>f</sup>The most common adverse events, occurring in at least 1% of the patients, that led to discontinuation of olaparib were nausea (2.1%), anemia (1.8%), fatigue (1.5%), and decreased neutrophil count (1.0%); there were no adverse events that occurred in at least 1% of patients that led to discontinuation of placebo.

<sup>g</sup>Adverse events leading to death are cardiac arrest (olaparib, n=1), AML (placebo, n=1), and ovarian cancer (placebo, n=1).

provided a statistically significant improvement in OS (HR 0.68; 98.5% CI 0.47-0.97; P = 0.009) with an absolute improvement in 4-year OS of 3.4% (89.8% olaparib; 86.4% placebo) in patients with high-risk EBC and gBRCA1/2pv following standardof-care chemotherapy, surgery, and radiation therapy, which if indicated had been completed at least 2 weeks before randomization. Updated descriptive analyses of IDFS and DDFS with the additional year of median follow-up demonstrated sustained absolute improvements (7.3% and 7.4%) for olaparib versus placebo in 4-year event-free rates, respectively. Safety analyses following completion of protocol therapy by all patients, including grade ≥3 AEs, SAEs, AEs leading to death, and AEs leading to discontinuation of treatment, demonstrated a favorable safety and tolerability profile consistent with the experience in the MBC setting with no substantive changes from the findings of the initial analysis. Although the key longterm safety endpoint of AML/MDS will require longer followup for complete assessment, the low incidence of 0.2% in the olaparib group and 0.3% in the placebo group with a median follow-up of 3.5 years coupled with the absence of new cases since the initial report is reassuring.

Breast cancers associated with gBRCA1/BRCA2pv are vulnerable to synthetic lethality caused by exposure to

PARP inhibitors that inhibit catalytic activities of PARP1 and trap PARP1 on DNA, creating lesions that require functional BRCA1 and BRCA2 protein for repair.<sup>3,4</sup> Because this vulnerability is independent of hormone receptor status, OlympiA was designed to assess the efficacy and safety of olaparib in patients with gBRCA1/2pv and high-risk, HER2negative EBC, irrespective of hormone receptor status. OlympiA was initially activated in patients with high-risk TNBC because of high unmet need for these patients in whom the residual recurrence risk following standard multimodality therapies remained sufficiently elevated to justify evaluating olaparib in the EBC setting, despite the lack of both phase III trial data and marketing authorization for olaparib in gBRCA1/2pv-associated MBC at that time. In contrast to gBRCA1pv-associated breast cancers, gBRCA2pvassociated breast cancers are predominantly hormone receptor positive.<sup>7,8</sup> Although adjuvant endocrine therapies reduce the risk of recurrence, patients presenting with larger, node-positive disease less responsive to NACT<sup>14,15</sup> or who have ≥4 positive axillary nodes at initial surgery have similar residual risk as patients with TNBC meeting eligibility criteria for OlympiA. Additionally, the complexities and challenges of conducting OlympiA made it unlikely that a new study specifically for patients with gBRCA1/2pv and hormone receptor-positive, high-risk EBC would be conducted. Therefore, once safety data on combinations of standard endocrine therapies and olaparib were available,<sup>17</sup> OlympiA was amended to include patients with hormone receptor-positive, HER2-negative EBC with risk of recurrence equivalent to the TNBC cohorts. Although the first patient with hormone receptor-positive disease was enrolled 18 months after start of accrual, the median follow-up was similar between the TNBC and hormone receptor-positive cohorts (3.6 versus 3.4 years).

Subgroup analyses of IDFS, DDFS, and OS demonstrate no evidence of heterogeneity for benefit of olaparib by hormone receptor status. The HR for olaparib relative to placebo for IDFS was 0.62 in TNBC (282 IDFS events in 1509 patients) and 0.68 in hormone receptor-positive disease (59 IDFS events in 325 patients), both less than the target HR of 0.7 for the ITT population (Figure 2B). The corresponding HR for DDFS was 0.59 (225 DDFS events) in the TNBC subgroup and 0.69 (54 DDFS events) in the hormone receptor-positive subgroup (Figure 2C). With relatively few deaths (n = 33) reported among the 325 patients with hormone receptor-positive EBC (Figure 2A), meaningful analysis of differential treatment effect on OS is highly constrained. Therefore, based on the negative test for heterogeneity by hormone receptor status and evidence for similar efficacy in IDFS and DDFS, coupled with the safety profile and the quality-of-life data, <sup>18</sup> patients with high-risk, hormone receptor-positive EBC should be considered for olaparib therapy. This conclusion is further supported by the lack of mechanistic rationale for differential synthetic lethal effects of PARP inhibition in a hormone receptor-positive context, evidence of similar treatment effect for PARP inhibitor therapy in MBC irrespective of hormone receptor status, 5,6 and reports of the randomized GeparOla study of olaparib in combination with paclitaxel, in which signals of comparative efficacy of olaparib/paclitaxel versus a carboplatin/paclitaxel regimen were stronger in the hormone receptor-positive subgroup. 19

OlympiA was notable for a relatively high adherence rate to study medication with 76% of the olaparib group completing at least 11 months of therapy compared with 82% of the placebo group. AEs were common reasons for discontinuation and the most common AEs leading to discontinuation were nausea and anemia. Nausea tends to occur early in treatment but diminishes in prevalence and grade with continued therapy. Patients should be informed of this potential side-effect and its likely time course and should be provided anti-emetic therapy to manage symptoms should they occur. Administering olaparib after a small meal may also help mitigate early nausea and potential vomiting.<sup>20</sup> Management of anemia on OlympiA included holding study medication until recovery of hemoglobin (Hb) to >9.5 g/dl. If recovery took >2 weeks, olaparib was reduced to 250 mg b.i.d. Study therapy was discontinued if repeated RBC transfusions were required to maintain the Hb >9.5 g/dl. This approach, adaptable to routine care, resulted in only 53 (5.8%) patients on olaparib requiring RBC transfusions compared with 8 (0.9%) on placebo (Supplementary Table S15A, available at https://doi.org/10. 1016/j.annonc.2022.09.159).

Following completion of accrual to OlympiA, KEYNOTE-522<sup>21</sup> demonstrated improved event-free survival (EFS) in TNBC with the addition of pembrolizumab to an NACT regimen of sequential carboplatin/paclitaxel followed by anthracycline with cyclophosphamide, followed by adjuvant pembrolizumab. Although the absolute improvement in EFS was 11% in patients without pathological complete response (pCR) with addition of pembrolizumab, 3-year EFS of this group was 67.4%, justifying consideration of additional post-surgical adjuvant therapy such as olaparib in patients with gBRCA1/2pv. Available safety data suggest that programmed cell death protein 1/programmed death-ligand 1 inhibitors can be co-administered with olaparib or other PARP1 inhibitors, 22,23 but this was not assessed in OlympiA.

The CREATE-X<sup>24</sup> study has also reported improvement in DFS (HR 0.58) and OS (HR 0.52) with adjuvant capecitabine in patients with TNBC and non-pCR following NACT that did not include platinum-based agents, which were allowed by OlympiA. A subsequent meta-analysis of 13 trials which evaluated capecitabine in EBC and included CREATE-X demonstrated improvement in DFS (HR 0.89) and OS (HR 0.83) in patients with TNBC.<sup>25</sup> There is an absence of safety data to support use of combination olaparib and capecitabine, so physicians and patients will need to choose between the two agents in the adjuvant setting. Although no data in EBC exist to inform the choice between the two agents, the OlympiAD MBC study in patients with gBRCA1/2pv demonstrated superiority of olaparib relative to mono-chemotherapy of physician's choice, in which the most common choice was capecitabine. 5 Similar findings were reported with talazoparib in the EMBRACA trial.<sup>6</sup> Additionally, there is evidence that patients with the basal subtype of TNBC may derive less benefit from capecitabine than their non-basal subtype affected counterparts, and patients with gBRCA1/2pv typically develop the basal subtype of TNBC. The most direct evidence comes from the GEICAM/ CIBOMA<sup>26</sup> open-label trial of adjuvant capecitabine following standard (N)ACT in early TNBC, stratified by basal versus nonbasal subtype based on immunohistochemistry staining for cytokeratin 5/6 and epidermal growth factor receptor. Although an HR of 0.82 (95% CI 0.63-1.06; P = 0.136) for the primary endpoint of DFS did not reach statistical significance, a pre-specified analysis by subtype suggested the smaller nonbasal cohort (26%) derived benefit from capecitabine with a DFS HR of 0.53 compared with an HR of 0.94 in the majority basal cohort. ECOG-ACRIN EA1131<sup>27</sup> was a randomized trial of adjuvant capecitabine versus platinum chemotherapy in patients with a basal subtype of TNBC determined by Prediction Analysis of Microarray 50 (PAM50) with  $\geq$ 1 cm of residual disease following taxane-based NACT. Accrual ended early when the IDMC determined that it was unlikely the study would demonstrate either noninferiority or superiority of platinum. Notably, 3-year IDFS in both arms was <50%, demonstrating high recurrence risks in this population despite use of either drug and the need for alternative approaches to mitigate this risk. These aggregate results, coupled with the favorable toxicity profile of olaparib in OlympiA, support the choice of olaparib in TNBC patients with gBRCA1/2pv.

Adjuvant therapy guidelines for high-risk, hormone receptor-positive breast cancer have been recently impacted by the monarchE trial, which demonstrated that 2 years of abemaciclib, co-administered with ET, improved 3year IDFS from 83.4% to 88.8% (HR 0.70; 95% CI 0.59-0.82).<sup>28</sup> There is an absence of safety data to support the use of a combination of olaparib, abemaciclib, and ET, so physicians and patients will need to choose between which of the two agents to combine with adjuvant ET. The monarchE trial has yet to demonstrate an improvement in OS and was not designed to assess the activity in patients with gBRCA1/2pv. Additionally, an evolving body of evidence suggests that patients with gBRCA2pv and hormone receptor-positive MBC may not respond as well to cyclindependent kinase 4 and 6 inhibitors. 29-31

In OlympiA, there was no evidence of statistical heterogeneity in the treatment effect for olaparib by hormone receptor status, and the similar HR for IDFS and DDFS for both hormone receptor-negative and hormone receptor-positive cohorts is consistent with a receptor-agnostic synthetic lethal targeting mechanism. The safety profile and quality-oflife data<sup>18</sup> from OlympiA also provide support that patients with gBRCA1/2pv and high recurrence risk, hormone receptor-positive EBC should be considered for combination adjuvant ET plus olaparib therapy following (N)ACT.

The pre-specified second IA of OlympiA with a median follow-up of 3.5 years demonstrates a statistically significant

improvement in OS with olaparib compared to placebo and maintenance of clinically meaningful absolute improvements in the previously reported statistically significant primary endpoint of IDFS and the secondary endpoint of DDFS. Subgroup analyses for all three endpoints demonstrate benefit irrespective of hormone receptor status, NACT versus ACT, prior use of platinum for breast cancer, and type of gBRCApv with CIs that include the point estimate of the HR in the overall population for each of the endpoints. The safety and tolerability profile of olaparib in this study remains consistent with that observed in previous studies of olaparib and only two cases (0.2%) of AML/MDS have been reported in the olaparib group compared with three (0.3%) in the placebo group. The results highlight the importance of testing for gBRCA1/2pv in patients with newly diagnosed high-risk EBC. Blinded follow-up of patients continues to assess long-term effects on risks for recurrent breast cancer and other second malignancies including AML/MDS, as well as to fully inform future translational studies to understand mechanisms of resistance to adjuvant olaparib.

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#### **DISCLOSURES**

CEG—uncompensated advisory board member for Astra-Zeneca, Genentech/Roche, Daiichi Sankyo, SeaGen and as compensated advisory board member for Exact Sciences. Medical writing assistance on manuscripts from Genentech/ Roche and Abbvie. Research funding from AstraZeneca, Genentech/Roche, Abbvie, Daiichi/Sankyo to institutions. Compensation for steering committee service to NSABP Foundation from Genentech/Roche. Accommodations and travel expenses from AstraZeneca, Genentech/Roche, Daiichi/Sankyo. RG-institutions have received research funding from AstraZeneca, MSD, Roche, and Novartis. MT—AstraZeneca employee and shareholder. LR—received salary support for project-related work under Agreement with Study Sponsor. PR-reports travel and accommodations by AZ. KC—employed by AstraZeneca and owns stock from AstraZeneca and BMS. AA—reports funding received by her institution as research funding from AstraZeneca, Roche/Genentech, Tesaro, Novartis, Pfizer, SERVIER, Biovica, GlaxoSmithKline, and Sanofi/Aventis, and royalties from Agendia for MammaPrint, due to the collaboration on the conduct of the MINDACT trial. GA—an employee of Merck Sharp & Dohme LLC, a subsidiary of Merck & Co., Inc., Rahway, NJ, USA and shareholder of Merck & Co., Inc., Rahway, NJ, USA. AA—advisory board MSD and Gilead; conference fees MSD and Gilead; spousal shares AstraZeneca; institutional research funding from Astra Zeneca. MA—(during the past 3 years) has received research funding (to Gustave Roussy) from Astra Zeneca and Eli-Lilly. Received honoraria (to Gustave Roussy): Astra Zeneca (to Institut Bergonié) Astra Zeneca; honoraria (to herself): Pfizer, Gilead. Travel grant: Astra Zeneca, Daiichi Sankyo. JBa-consulting fees from Astra Zeneca and Pfizer; travel expenses by Lilly; European patent request submitted (EP17382884.9) not related to this work. JBe—has received grants from Amgen, Astra Zeneca, Bayer, Merck KGaA, Pfizer, Roche, and Sanofi-Aventis to Karolinska Institutet and/or University Hospital. No personal payments. He also receives payment from UpToDate to Asklepios Medicine HB for a chapter on breast cancer prediction. He has recently been appointed board member of Wnt Research AB. JBI—(during the past 3 years) received research funding directly to the Institute of Cancer Research from: AstraZeneca, Merck KGaA, Puma Biotechnology, Pfizer, Roche, Novartis (previously GSK), Eli Lilly, Janssen-Cilag, and Clovis Oncology. SD—institutional grant from AstraZeneca during the conduct of this study; grants, all to her institution and all outside of the submitted work, from: Sanofi, Novartis, Lilly, Puma, Myriad, Orion, Amgen, Sanofi, Genomic Health, GE, Servier, Merck KGaA, BMS, Pierre Fabre, SeaGen, Exact Sciences, Rappta, Besins, Taiho, the European Commission, the French government, Foundation ARC; and non-financial support from Pfizer; AstraZeneca, and Roche Genentech. SMD—received research funding directly to the University of Pennsylvania from AstraZeneca and has received honoraria from AstraZeneca. AE—has received research funding directly to her institution from

AbbVie. **RNA** Astra Zeneca. and Diagnostics. FE—AstraZeneca employee and shareholder. LF—consulting/advisory role: Novartis, Pfizer, AstraZeneca/MSD; research funding to institution: AstraZeneca, MSD Oncology, Novartis. AF—AstraZeneca employee and AstraZeneca stockholder. JMF—has received institutional research grants from AstraZeneca, PUMA, Pfizer, Merus, Incyte, and Genentech. SF-reports support to her organization from AbbVie, AstraZeneca, Daiichi Sanko, Genentech, GlaxoSmithKline, MSD, and SeaGen. KG—advisory boards: AstraZeneca; Pfizer, Novartis, Lilly, MSD, Roche, SeaGen, Gilead, Ayala. Research funding: AstraZeneca, Pfizer, Roche, BMS. Speaker: Pfizer, Novartis, Lilly, AstraZeneca. LG—has served as compensated advisory board member for Astra-Zeneca, Daiichi Sankyo, and SeaGen. MG—reports personal fees/travel support from Amgen, DaiichiSankyo, AstraZeneca, EliLilly, LifeBrain, Veracyte, Novartis, PierreFabre, Merck KGaA; an immediate family member is employed by Sandoz. SH—AZ employee and shareholder. SAI—has advisory role for AstraZeneca, Bertis, Daiichi-Sankyo, Eisai, Eli Lilly, Hanmi, Idience, MSD, Novartis, Roche, Pfizer. Reports research grants from AstraZeneca, Boryung, Daewoong Pharm, Eisai, Roche, and Pfizer. SRL-institution has received honoraria for my role on the AstraZeneca International Breast Cancer Biomarker Advisory Board 2022. WJ—research grants and/or honoraria from: AstraZeneca, Celgene, Chugai, Daiichi/Sankyo, Eisai, ExactScience, GSK, Janssen, Lilly, Menarini, Merck KGaA, Novartis, Sanofi-Aventis, Roche, Pfizer, Seagen. BL-advisory boards for AZ, Pfizer, Merck KGaA, Lilly, Daiichi Sankyo, Gilead, SeaGen, and Novartis. SL—grants and honorarium to her institution from AbbVie, Amgen, AstraZeneca, Celgene, Daiichi Sankyo, Gilead, Novartis, Pfizer, Roche; and honorarium to her institution from Bristol Myers Squibb, Eli Lilly, Eirgenix, GlaxoSmithKline, Merck KGaA, Pierre Fabre, PriME/Medscape, and Seagen; grants to her institution from Cepheid; non-financial support for medical writing from Amgen, AstraZeneca, Bristol Myers Squibb, Celgene, Daiichi Sankyo, Gilead, Novartis, Pfizer, and Roche; patents EP21152186.9, EP18209672; EP15702464.7; and EP19808852.8 pending; and licensing fees from VM Scope GmbH, all to institution; personal fees from Chugai; personal employment with GBG Forschungs GmbH; personal non-financial interest in Gilead, Novartis, Pfizer, Roche, and SeaGen (steering committees); non-financial interests from AGO Member (German Gynaeological Oncology Society), ASCO Member, DKG Member (German Cancer Society) and ESMO (Member, Chair ESMO Breast (2019-21 and Steering Committee). PCL—reports stock ownership in Amgen, speaker honorarium from Schrodinger Inc., and unreimbursed consulting for BlueSphere Bio. FM-reports other from GBG research GmbH, during the conduct of the study; personal fees from Roche, AstraZeneca, Pfizer, Tesaro, Novartis, Amgen, PharmaMar, Genomic Health, CureVac, EISAI, Clovis, Janssen-Cilag, Gilead/Immunomedics, GSK, Merck KGaA, SeaGen, Myriad, and Pierre-Fabre, outside the submitted work. RM—receives salary support for project related work under

agreement with sponsors AstraZeneca, Roche, & GSK. KAP—has served as an uncompensated advisory board member for AstraZeneca. MP—scientific board member: Oncolytics; consultant/honoraria: AstraZeneca, Camel-IDS/ Precirix, Gilead, Immunomedics, Lilly, Menarini, MSD, Novartis, Odonate, Pfizer, Roche-Genentech, Seattle Genetics, Immutep, Seagen, NBE Therapeutics, Frame Therapeutics; research grants to institute: AstraZeneca, Immunomedics, Lilly, Menarini, MSD, Novartis, Pfizer, Radius, Roche-Genentech, Servier, Synthon. GR-reports funding received by her institution as research funding from AstraZeneca, Roche/Genentech, Tesaro, Novartis, Pfizer, SERVIER, Biovica, GlaxoSmithKline, and Sanofi/Aventis, and royalties received by her institution from Agendia for MammaPrint, due to the collaboration on the conduct of the MINDACT trial. RS—grants/contracts: AstraZeneca; participation on an advisory board: AstraZeneca, MSD, Clovis Oncology. ES-reports honoraria: Amgen, AstraZeneca, Cancérodigest, Clinigen, Curio Science, Egis, Eli Lilly, Exact Sciences, Gilead, high5md, Novartis, Oncompass Medicine, Pfizer, Pierre Fabre, Roche, Sandoz, TLC Biopharmaceuticals; travel support: Amgen, AstraZeneca, Egis, Gilead, Novartis, Pfizer, Roche; clinical research: Amgen, AstraZeneca, Eli Lilly, Novartis, Pfizer, Roche, Samsung; stock: AstraZeneca, Eli Lilly, Pfizer. PS-has received consulting fee and/or honoraria from Pfizer, MSD, Gilead, Seattle Genetics, Novartis, AstraZeneca, GSK, and research support (to the institution) from Novartis, Bristol-Meyers Squibb, MSD, Gilead. CFS—travel grants and speakers' honoraria from: Novartis, Roche, AstraZeneca, Gilead Sciences, SeaGen; research grants: AstraZeneca, Novartis, Amgen, and Daiichi-Sanyko. TŠ-patient advisory board honoraria: MSD, Pfizer, Gilead Sciences. ES—honoraria: Roche, Lilly, Pfizer, Merck KGaA, AstraZeneca, Novartis, SeaGen, Daiichi Sankyo/AstraZeneca, Gilead Sciences. Consulting or advisory role: Novartis, Roche, Pfizer, Lilly, AstraZeneca, Merck KGaA. Travel, accommodations, expenses: Novartis, Roche, Pfizer, Gilead Sciences. Uncompensated relationships: German Breast Group, MT—has received honoraria for lectures or chairs from Chugai, Takeda, Pfizer, Kyowa-Kirin, Taiho, Eisai, Daiichi-Sankyo, AstraZeneca, Eli Lilly, MSD, Exact Science, Novartis, Shimadzu, Yakult, Nippon Kayaku, and Devicore Medical Japan. He has served as compensated advisory board for Kyowa-Kirin, Daiichi-Sankyo, Eli Lilly, BMS, Athenex Oncology, Bertis, Terumo, and Kansai Medical Net. His institution has received research funding from Chugai, Takeda, Pfizer, Kyowa-Kirin, Taiho, JBCRG assoc., KBCRN assoc., Eisai, Eli Lilly, Daiichi-Sankyo, AstraZeneca, Astellas, Shimadzu, Yakult, Nippon Kayaku, AFI technology, Luxonus, Shionogi, GL Science, and Sanwa Shurui. He has served as uncompensated member of the board of directors for the association of JBCRG, the association of KBCRN, and the NPO organization OOTR. TAT—has consulting honoraria from advisory boards and research support from AstraZeneca. Also, honoraria from Pfizer. GV—received honoraria and consulting fees from Roche, AstraZeneca, Daiichi Sankyo, Merck KGaA,

Agilent, and Pfizer. YHP—reports grants from Roche, AstraZeneca, Pfizer, Novartis, and MSD; personal fees from AstraZeneca, Daiichi Sankyo, Eisai, Pfizer, MSD, Bixink, and Roche; and nonfinancial support from Pfizer and Hanmi. RY—reports consulting/advisor/honoraria: AstraZeneca, Eli Lilly, Gilead, Medison, MSD, Novartis, Pfizer, Roche, Teva; research grant: Roche. Research support: Exact Sciences. KHJ-consultancies (personal fees) from AstraZeneca, Bixink, Everest Medicine, MSD, Novartis, Pfizer, Roche, Takeda Pharmaceuticals. GSS—reports research support paid to the institution from Agendia, AstraZeneca, MSD, Roche, and SeaGen and consultancy fees paid to the institution from Biovica and SeaGen. MP—honoria/advisory board/educational events: Roche, Novartis, Pfizer, Eli Lilly, Exact Sciences, Veracyte, Pierre Fabre. MAC—reports research grant from Roche. He has served as Co-Chair of the Scientific Committee of IBCSG. MS—has received personal fees from AstraZeneca, BioNTech, Daiichi Sankyo, Eisai, Lilly, MSD, Novartis, Pantarhei Bioscience, Pfizer, Pierre Fabre, Roche, and SeaGen. His institution has received research funding from AstraZeneca, BioNTech, Eisai, Genentech, German Breast Group, Novartis, Palleos, Pantarhei Bioscience, Pierre Fabre, and SeaGen. In addition, he has a patent for EP 2390370 B1 and a patent for EP 2951317 B1 issued. AMB—reports consulting/advisor/honoraria: Roche, Celgene, AstraZeneca, Seagen, Daiichi Sankyo, Athenex, Lilly, MSD, Gilead, Novartis, Eisai, Pfizer, Samsung, Lilly, GE, Coherus, Puma; research funding to the institution: Roche, AstraZeneca, MSD, Novartis, Lilly, Gilead, Puma; travel, accommodation, expenses: Pfizer, Puma, GE. DC-Aptitude Health, Roche Sweden, Pfizer Limited, Celldex Therapeutics Inc, Carnall Farrar, ELI LILLY & Company, Astra Zeneca, Roche Products Ltd, Novartis Pharma AG, Novartis Pharmaceuticals Corporation, Pfizer Limited, PFS Ltd, Novartis Pharmaceuticals UK Limited, Merck KGaA, F. Hoffmann-La Roche AG, Clovis Oncology, Daiichi Sankyo, USA, Eisai, Exact Therapeutics, G1 Therapeutics, Galapagos NV, Genentech Inc, GSK (Glaxo SmithKline), Synthon Biopharmaceuticals BV: note name change to Byondis April 2020, Seagen, SANOFI, Sapience Therapeutics Ltd, Bexon/ Zymeworks Biopharmaceuticals Inc., NexGen, IQVIA. CC—received salary support for project-related work under Agreement with Study Sponsor. ANJT—reports consulting/ advisor/honoraria: Pfizer, Artios, Prime Oncology, Gilead, Merck KGaA; advisory board funds to institution: Gilead, AstraZeneca; research funding to the institution: AstraZeneca, Merck KGaA; expert testimony: EM Partners; stocks: Inbiomotion. Royalty associated payments—ICR rewards to inventor's scheme payments associated with patent for the use of PARP inhibitors in DNA deficient cancers, licensee--AstraZeneca. Other, grant funded by Breast Cancer Now (BCN) and Cancer Research UK (CRUK) to study homologous recombination deficient breast and other cancers, BCN/ CRUK receive payments associated with a patent for the use of PARP inhibitors in DNA deficient cancers, licensee-—AstraZeneca. All other authors have declared no conflicts of interest.

#### PREVIOUS RELATED WORKS

Tutt ANJ, Garber J, Gelber RD, et al. Pre-specified event driven analysis of overall survival (OS) in the OlympiA phase III trial of adjuvant olaparib (OL) in germline BRCA1/2 mutation (gBRCAm) associated breast cancer. ESMO 2022. Abstract VP1-March 2022.

Ganz PA, Bandos H, Spanic T, et al. Quality of life results from OlympiA: a phase III, multicenter, randomized, placebo-controlled trial of adjuvant olaparib after (neo)-adjuvant chemotherapy in patients with germline BRCA1/2 mutations and high-risk HER-2 negative early breast cancer (NSABP B-55). Presented at SABCS. December 10, 2021. Program Number: GS4-079 (Oral Abstract).

Tutt A, Garber JE, Kaufman B, et al. OlympiA: a phase III, multicenter, randomized, placebo-controlled trial of adjuvant olaparib after (neo) adjuvant chemotherapy in patients with germline BRCA1/2 mutations and high-risk HER2-negative early breast cancer. *J Clin Oncol.* 2021;39:18s (suppl; Abstract LBA1 ASCO Plenary).

Tutt ANJ, Garber JE, Kaufman B, et al. Adjuvant olaparib for patients with BRCA1- or BRCA2-mutated breast cancer. N Engl J Med. 2021;384(25):2394-2405.

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## APPENDIX 1. COLLABORATORS (PARTICIPATING GROUPS, ACCRUING INSTITUTIONS, AND LEAD INVESTIGATORS)

#### ABCSG: Austrian Breast and Colorectal Cancer Study Group

| Krankenhaus Hietzing, Abt. für Gynäkologie und Geburtshilfe         | Austria | Paul Sevelda                |
|---|---------|-----------------------------|
| KH Voecklabruck, Abt. f. Innere Medizin                             | Austria | Ferdinand<br>Haslbauer      |
| Krankenhaus der Barmherzigen Schwestern Ried                        | Austria | Monika Penzinger            |
| St. Josef KH, Interne Abt.  | Austria | Leopold Öhler               |
| LKH Leoben  | Austria | Christoph<br>Tinchon        |
| Universitätsklinikum Salzburg                                       | Austria | Richard Greil               |
| Klinikum Wels-Grieskirchen  | Austria | Sonja Heibl                 |
| Medizinische Universität Wien, Univ. Klinik für<br>Innere Medizin I | Austria | Rupert Bartsch              |
| Aerztezentrum—Ordination Dr. Viktor Wette                           | Austria | Viktor Wette                |
| Medizinische Universität Wien, Univ.Klinik für<br>Frauenheilkunde   | Austria | Christian F. Singer         |
| LKH Villach, Gynaekologisch-Geburtshilfliche Abt.                   | Austria | Claudia Pasterk             |
| Krankenhaus der Barmherzigen Schwestern Linz                        | Austria | Ruth Helfgott               |
| LKH-Universitätsklinikum Klinikum Graz                              | Austria | Gunda Pristauz-<br>Telsnigg |
| LKH-Universitätsklinikum Klinikum Graz                              | Austria | Herbert Stöger              |
| Elisabethinen Hospital  | Austria | Angsar<br>Weltermann        |
| Universitätsklinik Innsbruck  | Austria | Daniel Egle                 |
| Ordination Dr. Irene Thiel  | Austria | Irene Thiel                 |
| TumorZentrum Kepler Universitatsklinikum Linz                       | Austria | David Fuchs                 |
| LKH Rankweil  | Austria | Holger Rumpold              |
| Wilhelminenspital der Stadt Wien, 3. Med.<br>Abteilung              | Austria | Kathrin Strasser-<br>Weippl |

## AGO-B: Arbeitsgemeinschaft Gynäkologische Onkologie Breast Study Group

| Universitätsklinikum Freiburg                         | Germany Beate Rautenberg      |
|---|-------------------------------|
| Universitäts Hamburg-Eppendorf                        | Germany Volkmar Müller        |
| Universitätsmedizin Mainz                             | Germany Marcus Schmidt        |
| Klinikum rechts der Isar der TU Muenchen              | Germany Stefan Paepke         |
| Klinikum Bremen-Mitte                                 | Germany Mustafa Aydogdu       |
| Martin-Luther-Universität Halle-Wittenberg            | Germany Christoph<br>Thomssen |
| Klinikum Frankfurt Höchst GmbH                        | Germany Joachim Rom           |
| Helios-Kliniken Berlin—Buch                           | Germany Christine Mau         |
| Friedrich-Alexander-Universität Erlangen-<br>Nürnberg | Germany Peter Fasching        |
| Johanniter-Krankenhaus Bonn                           | Campany Huga Jackan           |
| Jonanniter-Krankennaus Bonn                           | Germany Uwe-Jochen<br>Göhring |
| Klinikum Esslingen GmbH                               | Germany Thorsten Kühn         |
| Gynäkologisch-onkologische Praxis                     | Germany Stefanie Noeding      |
| Universitätsklinikum Essen (AöR)                      | Germany Sherko Kümmel         |
| Marien Hospital Witten gGmbH                          | Germany John Hackmann         |
| Universitätsklinikum Aachen                           | Germany Elmar Stickeler       |

## BCT-ANZ: Breast Cancer Trials—Australia and New Zealand

| The Townsville Hospital       | Australia Abhishek Joshi      |
|-------------------------------|-------------------------------|
| Sir Charles Gairdner Hospital | Australia Joanna Dewar        |
| Prince of Wales Hospital      | Australia Michael Friedlander |
| Peter MacCallum Cancer Centre | Australia Kelly-Anne Phillips |
| Cabrini Hospital              | Australia Yoland Antill       |
| Mater Cancer Care Centre      | Australia Natasha Woodward    |
|                               | Continued                     |

| Appendix Continued                          |                                       |
|---|---------------------------------------|
| The Tweed Hospital                          | Australia Ehtesham Abdi               |
| Gosford Hospital                            | Australia Susan Tiley                 |
| Tamworth Rural Referral Hospital            | Australia Mathew George               |
| Royal Hobart Hospital                       | Australia David Boadle                |
| Concord Repatriation General Hospital       | Australia Annabel Goodwin             |
| Calvary Mater Newcastle                     | Australia Andre van der<br>Westhuizen |
| Ballarat Oncology & Haematology<br>Services | Australia George Kannourakis          |
| Royal Adelaide Hospital                     | Australia Nicholas Murray             |
| ICON Cancer Care Wesley                     | Australia Nicole McCarthy             |

#### **BOOG:** Borstkanker Onderzoek Groep

| Leids Universitair Medisch<br>Centrum                            | The Netherlands | Judith Kroep          |
|--|-----------------|-----------------------|
| Maastricht Universitair Medisch<br>Centrum                       | The Netherlands | Maaike de Boer        |
| Amphia Ziekenhuis  | The Netherlands | Joan Heijns           |
| Dutch Breast Cancer Research<br>Group, Utrecht, The Netherlands  | The Netherlands | Agnes Jager           |
| Zuyderland Medisch Centrum<br>Sittard-Geleen                     | The Netherlands | Franciscus<br>Erdkamp |
| Zaans Medisch Centrum  | The Netherlands | Sandra Bakker         |
| Nederlands Kanker Instituut<br>Antoni van Leeuwenhoek Ziekenhuis | The Netherlands | Gabe S. Sonke         |

#### CCTG: Canadian Cancer Trials Group

| Saskatchewan Cancer Agency                     | Canada | Amer Sami          |
|--|--------|--------------------|
| Cross Cancer Institute                         | Canada | John Mackey        |
| CISSSMC—Hospital Charles Le Moyne              | Canada | Catherine Prady    |
| Odette Cancer Centre, University of Toronto    | Canada | Andrea Eisen       |
| CHAUQ Hopital du St-Sacrement                  | Canada | Christine Desbiens |
| Centre Hospitalier de l'Universite de Montreal | Canada | Erica Patocskai    |
| Hopital General Juif                           | Canada | Cristiano Ferrario |
| BC—Vancouver Centre                            | Canada | Karen Gelmon       |
| Juravinski Cancer Centre                       | Canada | Louise Bordeleau   |
| Allan Blair Cancer Centre                      | Canada | Haji Chalchal      |
| CancerCare Manitoba                            | Canada | Saroj Niraula      |

#### CEEOG: Central and East European Oncology Group

| Tel Aviv Sourasky Medical Center Ichilov   | Israel | Ido Wolf        |
|--|--------|-----------------|
| Uniwersyteckie Centrum Kliniczne w Gdańsku | Poland | Elżbieta Senkus |

# **EORTC:** European Organisation for Research and Treatment of Cancer

| Cliniques Universitaires Saint-Luc                      | Belgium | François Duhoux                |
|---|---------|--------------------------------|
| A.Z. Damiaan  | Belgium | Randal d'Hondt                 |
| Cliniques universitaires de<br>Bruxelles—Hôpital Érasme | Belgium | Sylvie Luce                    |
| Institut Jules Bordet                                   | Belgium | Daphné t'Kint de<br>Roodenbeke |
| Universitair Ziekenhuis Antwerpen (UZA)                 | Belgium | Konstantinos<br>Papadimitriou  |
| AZ Groeninge  | Belgium | Marleen Borms                  |
| CHU UCL Namur   | Belgium | Claire Quaghebeur              |
| Institut du Cancer de Montpellier Val<br>d'Aurelle      | France  | William Jacot                  |
| Institut Curie—Hôpital René Huguenin                    | France  | Etienne Brain                  |
|   |         | Continued                      |

| Appendix Continued  |          |                           |
|---|----------|---------------------------|
| CHU de Limoges—Hôpital Dupuytren                            | France   | Laurence Venat-<br>Bouvet |
| Hôpital Privé du Confluent                                  | France   | Alain Lortholary          |
| Narodowy Instytut Onkologii im. Marii<br>Skłodowskiej-Curie | Poland   | Zbigniew Nowecki          |
| Centro Clínico Champalimaud                                 | Portugal | Fátima Cardoso            |
| Western General Hospital                                    | UK       | Richard Hayward           |

# GAICO: Grupo Argentino De Investigación Clinica En Oncologia

| Clinica Universitaria Privada Reina Fabiola                  | Argentina | Santiago Bella                |
|--|-----------|-------------------------------|
| Centro Oncologico de Integracion Regional                    | Argentina | Mauricio Fernández<br>Lazzaro |
| Clínica Privada Colombo                                      | Argentina | Norma Pilnik                  |
| Instituto de Oncología de Rosario                            | Argentina | Luis E. Fein                  |
| Clinica ISIS   | Argentina | Cesar Blajman                 |
| CENIT Centro Medico de Neuro,<br>Investigacion y Tratamiento | Argentina | Guillermo Lerzo               |
| Centro de Oncologia e Investigacion en<br>Buenos Aires       | Argentina | Mirta Varela                  |
| Centro Medico San Roque                                      | Argentina | Juan Jose Zarba               |
| Centro Oncologico Riojano Integral (Cori)                    | Argentina | Diego Kaen                    |
| Instituto Medico Especializado Alexander                     | Argentina | Maria Victoria                |
| Fleming  |           | Constanzo                     |

## GBG: German Breast Group

| Universitätsklinikum Münster                              | Germany | Joke Tio                     |
|---|---------|------------------------------|
| Henriettenstiftung, Hannover                              | Germany | Wulf Siggelkow               |
| Klinikum Offenbach  | Germany | Christian Jackisch           |
| Klinikum der Eberhard-Karls-Universität                   | Germany | Eva Maria Grischke           |
| Tübingen  |         |                              |
| Wald-Klinikum Gera  | Germany | Dirk Zahm                    |
| DONAUISAR Klinikum Deggendorf                             |         | Sara Tato-Varela             |
| Elisabeth-Krankenhaus Kassel                              | Germany | Sabine Schmatloch            |
| Praxisklinik Berlin                                       | Germany | Peter Klare                  |
| Johanniter-Krankenhaus der Altmark<br>Stendal             | Germany | Andrea Stefek                |
| Universitätsklinikum Köln                                 | Germany | Kerstin Rhiem                |
| Universitätsklinikum Essen (AöR)                          | Germany | Oliver Hoffmann              |
| Kliniken Essen-Mitte                                      | Germany | Sherko Kümmel                |
| Caritasklinik St. Theresia, Saarbrücken                   | Germany | Mustafa Deryal               |
| Praxis und Tagesklinik, Ebersberg                         | Germany | Isolde Gröll                 |
| Städtisches Klinikum Brandenburg                          | Germany | Peter Ledwon                 |
| Gemeinschaftspraxis, Hildesheim                           | Germany | Christoph Uleer              |
| Klinikum Chemnitz   | Germany | Petra Krabisch               |
| Ev. Waldkrankenhaus Spandau, Berlin                       | Germany | Jochem Potenberg             |
| Luisenkrankenhaus GmbH&Co.KG<br>Düsseldorf                | Germany | Maren Darsow                 |
| Medizinische Hochschule Hannover                          | Germany | Tjoung-Won Park-<br>Simon    |
| MVZ Osthessen GmbH, Fulda                                 | Germany | Heinz-Gert Höffkes           |
| Oncologianova GmbH, Recklinghausen                        | Germany | Till-Oliver Emde             |
| Studienzentrum Zehlendorf, Berlin                         | Germany | Gerd Graffunder              |
| StVincentius Kliniken gAG Karlsruhe                       | Germany | Oliver Tomé                  |
| Universitätsklinikum Leipzig AöR                          | Germany | Dirk Forstmeyer              |
| Praxis Dr. med. Jürgen Terhaag, Eggenfelden               | Germany | Jürgen Terhaag               |
| Rotkreuzklinikum Munich                                   | Germany | Christoph Salat              |
| Universitätsklinikum Carl Gustav Carus der<br>TU Dresden  | Germany | Karin Kast                   |
| Gemeinschaftspraxis für Hämatologie und Onkologie, Erfurt | Germany | Steffi Weniger               |
| Onkologisch Hämatologische<br>Schwerpunktpraxis, Bremen   | Germany | Carsten Schreiber            |
| Gemeinschaftspraxis, Augsburg                             | Germany | Bernhard Heinrich            |
| Klinikum Südstadt, Rostock                                | Germany | Max Dieterich                |
| St. Vincenz Krankenhaus, Karlsruhe                        | Germany | Michaela Penelope<br>Wüllner |

## **GEICAM: Spanish Breast Cancer Group**

| Hospital Clinico Universitario Lozano Blesa Hospital Clinico Universitario San Carlos Spain José Ángel García Sáenz Complejo Hospitalario Universitario A Coruña Consorci Sanitari de Terrassa Hospital Arnau de Vilanova (Lleida) Hospital Universitario Virgen Macarena Spain Serafín Morales Murillo Hospital Universitario Virgen Macarena Spain Fernando Henao Carrasco Spain Salvador Blanch Tormo Oncología (IVO) Hospital Universitario de Donostia Hospital Universitario de Donostia Spain Isabel Álvarez López Spain Juan Ignacio Delgado Mingorance Hospital Lucus Augusti de Lugo Spain Isabel Álvarez Gomez Clínica Universitaria de Navarra Spain Marta Santisteban Spain Josefina Cruz Jurado (Tenerife) Hospital Universitario de Canarias (Tenerife) Hospital Universitario Virgen del Rocio Hospital Universitario de Castello Spain Manuel Ruiz Borrego Spain Eduardo Martínez de Dueñas Complejo Asistencial de Avila Spain Jose Enrique Alés Martínez Dueñas Spain Juan De la Haba Hospital Universitario Reina Sofía Spain Juan De la Haba Hospital Universitario de Elche Hospital Miguel Servet Spain Alvaro Rodríguez Lescure Spain Alvaro Rodríguez Lescure Spain Alvaro Rodríguez Lescure Spain Antonio Antón Torres Spain Gema Llort Crusades Spain Santiago González-Santiago Spain Antonia Marquez Aragones Victoria Spain Alvaro Rodríguez Lescure Spain Antonia Marquez Aragones Victoria Spain Gema Llort Crusades Spain Agusti Barnadas Molins Spain José Ignacio Chacón López-Muñiz Spain Alvaro Rodríguez Spain Alvaro Rodríguez Aragones Victoria Complejo Hospitalario de Jaen Spain Antonia Marquez Aragones Spain Agusti Barnadas Molins Spain José Ignacio Chacón López-Muñiz Spain Alvaro Rodríguez |  |                                  |
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| Coruña Consorci Sanitari de Terrassa Consorci Sanitari de Terrassa Hospital Arnau de Vilanova (Lleida) Hospital Universitario Virgen Macarena Fundación Instituto Valenciano de Oncología (IVO) Hospital Universitario de Donostia Hospital Universitario de Donostia Spain Isabel Álvarez López Hospital Infanta Cristina Spain Isabel Álvarez López Hospital Infanta Cristina Spain Isabel Álvarez López Hospital Infanta Cristina Spain Juan Ignacio Delgado Mingorance Hospital Lucus Augusti de Lugo Clínica Universitaria de Navarra Hospital Universitario de Canarias (Tenerife) Hospital Germans Trias i Pujol Hospital Germans Trias i Pujol Hospital Universitario Virgen del Rocio Hospital Universitario Virgen del Rocio Hospital Universitario Reina Sofía Hospital Universitario Ramón y Cajal Hospital Universitario Ramón y Cajal Hospital General Universitario de Elche Hospital Miguel Servet Corporació Sanitària Parc Taulí Hospital San Pedro de Alcántara Formal Miguel Servet Corporació Sanitària Parc Taulí Hospital Clínico Univ. Virgen de la Victoria Complejo Hospitalario de Jaen Hospital de la Santa Creu i Sant Pau Toledo, H. V. de la Salud, Oncología Hospital Universitari i Politècnic La Fe Hospital Universitario de Salamanca  |  | ·                                |
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| Mingorance  Hospital Lucus Augusti de Lugo Clínica Universitaria de Navarra Hospital Universitario de Canarias (Tenerife) Hospital Germans Trias i Pujol Hospital Universitario Virgen del Rocio Hospital Provincial Centre de Castello Hospital Provincial Centre de Castello Hospital Universitario Reina Sofía Hospital Miguel Servet Corporació Sanitària Parc Taulí Hospital San Pedro de Alcántara Complejo Hospitalario de Jaen Hospital Clínico Univ. Virgen de la Victoria Complejo Hospitalario de Jaen Hospital de la Santa Creu i Sant Pau Toledo, H. V. de la Salud, Oncología Hospital Universitari i Politècnic La Fe Hospital Universitari i Politècnic La Fe Hospital Clínico Universitario de Salamanca Mingorance Spain Elena Álvarez Gomez Spain Anauel Ruiz Borrego Spain Manuel Ruiz Borrego Spain Antonia Martínez Jañez Spain Antonia Antorio Torres Spain Antonio Antón Torres Spain Antonia Marquez Aragones Victoria Complejo Hospitalario de Jaen Spain Ana Laura Ortega Spain Angusti Barnadas Molins Spain José Ignacio Chacón López-Muñiz Spain Miguel Martín Jiménez   | Hospital Universitario de Donostia       | Spain Isabel Álvarez López       |
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| Hospital Quiron de Madrid Spain Lucía González Cortijo   |  | Spain César Rodríguez            |
|  | Hospital Quiron de Madrid                | Spain Lucía González Cortijo     |

## GOIRC: Italian Oncology Group for Clinical Research

| Ospedale Generale Regionale Bolzano Boheler<br>Lorenz       | Italy Elisabetta<br>Cretella |
|---|------------------------------|
| Azienda Ospedaliera Policlinico di Modena                   | Italy Laura Cortesi          |
| Ospedale di Belcolle  | Italy Enzo Maria<br>Ruggeri  |
| AO Busto Arsizio—Presidio di Saronno—SC<br>Oncologia Medica | Italy Claudio Verusio        |
| Ospedale Sacro Cuore  | Italy Stefania Gori          |
| Azienda Ospedaliera "Mater Salutis"/Aulss 9                 | Italy Andrea Bonetti         |
| Ospedale S.Maria della Misericordia                         | Italy Anna Maria<br>Mosconi  |

## IBCG: Icelandic Breast Cancer Group

| Landsnitali | , University Hospital | Iceland | Oskar Johannsson |
|-------------|-----------------------|---------|------------------|
|             |                       |         |                  |

## IBCSG: International Breast Cancer Study Group

| CHU de Liège   | Belgium | Guy<br>Jerusalem |
|--|---------|------------------|
| UZ Leuven  | Belgium | Patrick Neven    |
| Országos Onkológiai Intézet Chemotherapy<br>Department "B" | Hungary | Tünde Nagy       |
|  |         | Continued        |

| Appendix Continued                            |             |                      |
|---|-------------|----------------------|
| A. O. Ospedale di Circolo e Fondazione MACCHI | Italy       | Graziella<br>Pinotti |
| European Institute of Oncology                | Italy       | Marco A.<br>Colleoni |
| Fondazione S. Maugeri                         | Italy       | Antonio<br>Bernardo  |
| Ospedale Infermi—Rimini, AUSL della Romagna   | Italy       | Lorenzo<br>Gianni    |
| Multimedica Castellanza                       | Italy       | Eraldo Bucci         |
| Ospedale Misericordia e Dolce                 | Italy       | Laura<br>Biganzoli   |
| University Hospital of Zurich                 | Switzerland | Konstantin<br>Dedes  |
| Inselspital Bern                              | Switzerland | Urban Novak          |
| Centre Hospitalier Universitaire Vaudois      | Switzerland | Khalil Zaman         |

# ICR CTSU: Institute of Cancer Research—Clinical Trials and Statistics Unit

| Bristol Royal Infirmary, Dept of Oncology       | UK Jeremy            |
|---|----------------------|
|   | Braybrooke           |
| Weston Park Hospital, Oncology                  | UK Matthew Winter    |
| Queen Elizabeth Hospital                        | UK Daniel Rea        |
| St Georges Hospital, Dept of Oncology           | UK Muireann Kelleher |
| The Beatson West of Scotland Cancer Centre      | UK Sophie Barrett    |
| Nottingham City Hospital                        | UK Stephen Chan      |
| Royal Bournemouth Hospital                      | UK Tamas Hickish     |
| Belfast City Hospital                           | UK Jane Hurwitz      |
| St Bartholomew's Hospital                       | UK John Conibear     |
| CNS/Manager for Cancer and Haematology Clinical | UK Apurna            |
| Trials  | Jegannathen          |
| Royal Marsden Hospital                          | UK Marina Parton     |
| Guys And St Thomas Hospital                     | UK Andrew Tutt       |
| Russells Hall Hospital                          | UK Rozenn Allerton   |
| Velindre Cancer Centre                          | UK Annabel Borley    |
| The Christie Hospital NHS Foundation Trust      | UK Anne Armstrong    |
| Southampton General Hospital                    | UK Ellen Copson      |
| Churchill Hospital                              | UK Nicola Levitt     |
| Addenbrooke's Hospital                          | UK Jean Abraham      |
| St James' University Hospital                   | UK Timothy Perren    |
| University College Hospitals London             | UK Rebecca Roylance  |

## JBCRG: Japan Breast Cancer Research Group

| Iwate Medical University Hospital              | Japan Kazushige Ishida |
|--|------------------------|
| Nagoya City University Hospital                | Japan Tatsuya Toyama   |
| National Hospital Organization Osaka National  | Japan Norikazu Masuda  |
| Hospital                                       |                        |
| Shizuoka Cancer Center                         | Japan Junichiro        |
|  | Watanabe               |
| National Hospital Organization Kyushu Cancer   | Japan Eriko Tokunaga   |
| Center   |                        |
| National Cancer Center Hospital                | Japan Takayuki         |
|  | Kinoshita              |
| Hakuaikai Sagara Hospital                      | Japan Yoshiaki Rai     |
| Kyoto University Hospital                      | Japan Masahiro Takada  |
| Gunma Prefectural Cancer Center                | Japan Yasuhiro         |
|  | Yanagita               |
| Chiba Cancer Center                            | Japan Rikiya Nakamura  |
| Osaka International Cancer Institute           | Japan Takahiro         |
|  | Nakayama               |
| Osaka University Hospital                      | Japan Yasuto Naoi      |
| Aichi Cancer Center Hospital                   | Japan Hiroji Iwata     |
| Showa University Hospital                      | Japan Seigo Nakamura   |
| National Hospital Organization Hokkaido Cancer | Japan Masato           |
| Center   | Takahashi              |
|  | Japan Kenjiro Aogi     |
|  | Continued              |

| Appendix Continued                                      |                        |
|---|------------------------|
| National Hospital Organization Shikoku Cancer<br>Center |                        |
| St Marianna University School of Medicine               | Japan Koichiro Tsugawa |
| National Cancer Center Hospital East                    | Japan Hirofumi Mukai   |
| The Cancer Institute Hospital of JFCR                   | Japan Toshimi Takano   |
| Saitama Medical University International Medical Center | Japan Akihiko Osaki    |
| Niigata Cancer Center Hospital                          | Japan Nobuaki Sato     |
| St. Luke's International Hospital                       | Japan Hideko Yamauchi  |
| Tokai University Hospital                               | Japan Yutaka Tokuda    |
| Hiroshima City Hospital                                 | Japan Mitsuya Ito      |
| Kochi Medical School Hospital                           | Japan Takeki Sugimoto  |

NCI National Clinical Trials Network: Comprised of NRG Oncology, Alliance for Clinical Trials in Oncology, ECOG-ACRIN Cancer Research Group and Southwest Oncology Group

| Banner MD Anderson Cancer Center  USA Shakeela W. Bahadur  UCLA/Jonsson Comprehensive Cancer Center  USC/Norris Comprehensive Cancer Center  USA Min J. Lu  Los Angeles County-USC Medical Center  City of Hope Comprehensive Cancer Center  Lisa Monica M. Mita  City of Hope Comprehensive Cancer Center  Kaiser Permanente—Fontana  Stanford Cancer Institute Palo Alto  Kaiser Permanente San Leandro  Kaiser Permanente—Northern California  Kaiser Permanente—Northern California  Kaiser Permanente—Northern California  Kaiser Permanente—Southern California  Kaiser Permanente—Southern California  Kaiser Permanente—Colorado  Kaiser Permanente—Santa Teresa-San Jose  Saint Joseph's Medical Center  Kaiser Permanente—Fresno  Sutter Cancer Research Consortium—Sacramento  Kaiser Permanente—Woodland Hills  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Shanda Hills  Kaiser |   |  |
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| UCLA/Jonsson Comprehensive Cancer Center USC/Norris Comprehensive Cancer Center USC Min J. Lu Los Angeles County-USC Medical Center USC Min J. Lu Cedars-Sinai Medical Center USC Monica M. Mita City of Hope Comprehensive Cancer Center USC James Waisman Kaiser Permanente—Fontana USC Jonathan A. Polikoff Stanford Cancer Institute Palo Alto USC Melinda L. Telli Kaiser Permanente—San Leandro USC Samantha A. Seaward Kaiser Permanente—Northern California USC Samantha A. Seaward Kaiser Permanente—Northern California USC James Waisman Kaiser Permanente—Northern California USC James Waisman USC James Waisman USC Samantha A. Seaward USC James Waisman Kaiser Permanente—Northern California USC James Waisman USC James Waisman USC James Waisman Kaiser Permanente—Southern California USC James Waisman USC Jame | Banner MD Anderson Cancer Center  |  |
| USC/Norris Comprehensive Cancer Center Los Angeles County-USC Medical Center Cedars-Sinai Medical Center City of Hope Comprehensive Cancer Center USA Min J. Lu USA Monica M. Mita City of Hope Comprehensive Cancer Center USA James Waisman Kaiser Permanente—Fontana USA Jonathan A. Polikoff Stanford Cancer Institute Palo Alto USA Melinda L. Telli Kaiser Permanente San Leandro USA Samantha A. Seaward Kaiser Permanente—Vallejo USA J. Marie Suga Kaiser Permanente—Northern California USA Samantha A. Seaward Kaiser Permanente Oncology Clinical Trials—Northern California USA J. Marie Suga USA J. Marie Suga Trials—Permanente—Southern California USA Jennifer Fu Carney Kaiser Permanente—Colorado USA Alex Menter Kaiser Permanente—Santa Teresa-San Jose USA J. Marie Suga Saint Joseph's Medical Center USA Ajithkumar Puthillath Kaiser Permanente—Fresno USA J. Marie Suga Sutter Cancer Research Consortium—Sacramento USA J. Marie Suga Sutter Cancer Research Consortium—Sacramento USA J. Marie Suga Kaiser Permanente—Woodland Hills USA Jonathan A. Polikoff Kaiser Permanente—Baldwin Park USA Jonathan A. Polikoff Contra Costa Regional Medical Center USA James H. Feusner Sutter Cancer Research Consortium—Roseville USA Jonathan A. Polikoff Contra Costa Regional Medical Center USA Jonathan A. Polikoff Contra Costa Regional Medical Center USA Jonathan A. Polikoff Contra Costa Regional Medical Center USA Jonathan A. Polikoff Contra Costa Regional Medical Center—Vacaville USA Jonathan A. Polikoff Discortium—Roseville USA Jonathan A. Polikoff Contra Costa Regional Medical Center—Vacaville USA Jonathan A. Polikoff Discortium—Roseville USA Jonathan A. Polikoff Contra Costa Regional Medical Center—Vacaville USA Jonathan A. Polikoff Discortium—Roseville USA Jonathan A. Polikoff  | LICIA/Isassas Communication Communication   |  |
| Los Angeles County-USC Medical Center Cedars-Sinai Medical Center City of Hope Comprehensive Cancer Center Kaiser Permanente—Fontana Stanford Cancer Institute Palo Alto Stanford Cancer Institute Palo Alto USA Melinda L. Telli USA Samantha A. Seaward Seaward Kaiser Permanente—Vallejo Staiser Permanente—Northern California Seaward Sea |   |  |
| Cedars-Sinai Medical Center City of Hope Comprehensive Cancer Center Kaiser Permanente—Fontana  Stanford Cancer Institute Palo Alto Kaiser Permanente San Leandro  Kaiser Permanente—Vallejo Kaiser Permanente—Northern California  Kaiser Permanente Oncology Clinical Trials—Northern California  Kaiser Permanente—Southern California  Kaiser Permanente—Sonta Teresa-San Jose  Saint Joseph's Medical Center  Kaiser Permanente—Santa Teresa-San Jose  Saint Joseph's Medical Center  WSA Ji Marie Suga  Saint Joseph's Medical Center  WSA Jonathan A. Polikoff  Kaiser Permanente—Fresno  Sutter Cancer Research Consortium—Sacramento  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Woodland Hills  WSA Jonathan A. Polikoff  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  WSA Jonathan A. Polikoff  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  WSA Jonathan A. Polikoff  Disenberg  Kaiser Permanente West Los Angeles  USA Jonathan A. Polikoff  Disenberg  Kaiser Permanente Medical Center—Vacaville  WSA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale  USA Derrick Wong  University of Colorado Cancer Center  USA Derrick Wong  University of Colorado Cancer Center  USA Peter D. Eisenberg  Kaiser Permanente Medical Center—Trumbull  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospi |   |  |
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| Kaiser Permanente—Fontana USA Jonathan A. Polikoff Stanford Cancer Institute Palo Alto Kaiser Permanente San Leandro Kaiser Permanente—Vallejo Kaiser Permanente—Northern California Kaiser Permanente Oncology Clinical Trials—Northern California USA J. Marie Suga USA Jennifer Fu Carney Kaiser Permanente—Southern California USA Jennifer Fu Carney Kaiser Permanente—Colorado USA Alex Menter USA Jennifer Fu Carney USA Ajithkumar Puthillath Kaiser Permanente—Santa Teresa-San Jose USA J. Marie Suga Saint Joseph's Medical Center USA Jonathan A. Polikoff Kaiser Permanente—Fresno USA J. Marie Suga USA Jonathan A. Polikoff USA Peter D. Eisenberg USA Jonathan A. Polikoff Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong University of Colorado Cancer Center USA Alexander T. Urquhart Yale University USA Erin W. Hofstatter USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull USA Erin W. Hofstatter   |   |  |
| Stanford Cancer Institute Palo Alto  Kaiser Permanente San Leandro  Kaiser Permanente—Vallejo  Kaiser Permanente—Vallejo  Kaiser Permanente—Northern California  Kaiser Permanente Oncology Clinical  Trials—Northern California  Kaiser Permanente—Southern California  Kaiser Permanente—Southern California  Kaiser Permanente—Hawaii  Kaiser Permanente—Colorado  Kaiser Permanente—Colorado  Kaiser Permanente—Santa Teresa-San Jose  Saint Joseph's Medical Center  Kaiser Permanente—Fresno  Sutter Cancer Research Consortium—Sacramento  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Woodland Hills  Kaiser Permanente—Baldwin Park  Kaiser Permanente—Baldwin Park  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  Kaiser Permanente West Los Angeles  Kaiser Permanente Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  Kaiser Permanente West Los Angeles  VISA Jonathan A.  Polikoff  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  VISA Jonathan A.  Polikoff  Marin Cancer Care Inc  Wisa Peter D.  Eisenberg  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente—San Marcos  USA Jonathan A.  Polikoff  Palo Alto Medical Foundation—Sunnyvale  UISA Derrick Wong  University of Colorado Cancer Center  USA Alexander T.  Urquhart  Yale University  VISA Erin W. Hofstatter  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer | , , ,   |  |
| Kaiser Permanente San Leandro  Kaiser Permanente—Vallejo  Kaiser Permanente—Northern California  Kaiser Permanente Oncology Clinical Trials—Northern California  Kaiser Permanente—Southern California  Kaiser Permanente—Southern California  Kaiser Permanente—Southern California  Kaiser Permanente—Hawaii  USA J. Marie Suga  USA J. Marie Suga  USA J. Marie Suga  USA Jennifer Fu Carney  Kaiser Permanente—Hawaii  USA Jennifer Fu Carney  Kaiser Permanente—Colorado  Kaiser Permanente—Santa Teresa-San Jose  Saint Joseph's Medical Center  Kaiser Permanente Los Angeles Medical Center  WSA Ajithkumar Puthillath  Kaiser Permanente—Fresno  Sutter Cancer Research Consortium—Sacramento  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Woodland Hills  WSA J. Marie Suga  Saint Joseph's Medical Center  USA J. Marie Suga  USA J. Marie Suga  USA J. Marie Suga  USA J. Marie Suga  WSA J. Marie Suga  WSA J. Marie Suga  WSA J. Marie Suga  WSA Jonathan A. Polikoff  Kaiser Permanente—Baldwin Park  USA Jonathan A. Polikoff  Kaiser Permanente—Baldwin Park  USA Janes H. Feusner  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  USA Jonathan A. Polikoff  Marin Cancer Care Inc  USA Peter D. Eisenberg  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente Medical Center—Vacaville  VSA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale  UNA Perrick Wong  UNA Porrick Wong  UNA Porrick Wong  UNA Porrick Wong  UNA Porrick Wong  UNA Polikoff  Was Perrick Was Perrick Was Perrick Was Perrick Was Perrick Wong  UNA Polikoff  Was Perrick Was Perrick Was Perrick Was P | Kaiser Permanente—Fontana   |  |
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| Kaiser Permanente—Vallejo Kaiser Permanente—Northern California Kaiser Permanente Oncology Clinical Trials—Northern California Kaiser Permanente—Southern California Kaiser Permanente—Southern California Kaiser Permanente—Southern California Kaiser Permanente—Hawaii  Kaiser Permanente—Hawaii  Kaiser Permanente—Colorado  Kaiser Permanente—Santa Teresa-San Jose Saint Joseph's Medical Center  Kaiser Permanente—Santa Teresa-San Jose Saint Joseph's Medical Center  WSA J. Marie Suga Saint Joseph's Medical Center  WSA J. Marie Suga Sutter Cancer Research Consortium—Sacramento Kaiser Permanente—Fresno Sutter Cancer Research Consortium—Sacramento Kaiser Permanente—Santa Rosa WSA J. Marie Suga Sutter Cancer Research Consortium—Sacramento WSA J. Marie Suga WSA Jonathan A. Polikoff  Contra Costa Regional Medical Center USA Jonathan A. Polikoff  Contra Costa Regional Medical Center WSA Jonathan A. Polikoff Marin Cancer Research Consortium—Roseville WSA Jonathan A. Polikoff Marin Cancer Care Inc WSA Jonathan A. Polikoff Marin Cancer Care Inc USA J. Marie Suga Waiser Permanente Medical Center—Vacaville WSA J. Marie Suga Waiser Permanente Medical Center—Vacaville WSA J. Marie Suga Waiser Permanente Medical Center—Vacaville WSA J. Marie Suga Waiser Permanente—San Marcos USA J. Marie Suga Waiser Permanente Medical Center—Vacaville WSA Peter D. Eisenberg Waisen Peter San Waisen Peter Waisen Waisen Peter Waisen  | Kaiser Permanente San Leandro   |  |
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| Kaiser Permanente Oncology Clinical Trials—Northern California Kaiser Permanente—Southern California Kaiser Permanente—Southern California  Kaiser Permanente—Hawaii  Kaiser Permanente—Hawaii  USA Jara N. Durna  USA Jera N. Durna  USA Jernifer Fu Carney  Kaiser Permanente—Colorado  Kaiser Permanente—Santa Teresa-San Jose USA J. Marie Suga  Saint Joseph's Medical Center  VSA Ajithkumar Puthillath  Kaiser Permanente Los Angeles Medical Center  USA Ajithkumar Puthillath  Kaiser Permanente—Fresno USA J. Marie Suga  Sutter Cancer Research Consortium—Sacramento USA J. Marie Suga  Kaiser Permanente—Santa Rosa USA J. Marie Suga  Kaiser Permanente—Woodland Hills USA Jonathan A. Polikoff  Kaiser Permanente—Baldwin Park USA Jonathan A. Polikoff  Contra Costa Regional Medical Center USA James H. Feusner  Sutter Cancer Research Consortium—Roseville USA Kristie A Bobolis  Kaiser Permanente West Los Angeles USA J. Marie Suga  Kaiser Permanente Medical Center—Vacaville USA Pelter D. Eisenberg  Kaiser Permanente Medical Center—Vacaville USA J. Marie Suga  Kaiser Permanente Medical Center—Vacaville USA Pelter D. Eisenberg  Kaiser Permanente—San Marcos USA J. Marie Suga  Kaiser Permanente—San Marcos USA Polikoff  Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong  University of Colorado Cancer Center USA Alexander T. Urquhart  Yale University Smilow Cancer Hospital-Waterbury Care Center USA Erin W. Hofstatter  Smilow Cancer Hospital-Waterbury Care Center USA Erin W. Hofstatter  Medstar Franklin Square Medical Center/Weinberg USA Edward C. McCarron   | •   |  |
| Trials—Northern California  Kaiser Permanente—Southern California  Kaiser Permanente—Hawaii  Kaiser Permanente—Hawaii  USA Jennifer Fu Carney  Kaiser Permanente—Santa Teresa-San Jose  Saint Joseph's Medical Center  Saint Joseph's Medical Center  Saint Joseph's Medical Center  WSA Ajithkumar Puthillath  Kaiser Permanente—Fresno  Suther Cancer Research Consortium—Sacramento  Kaiser Permanente—Fresno  Sutter Cancer Research Consortium—Sacramento  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Woodland Hills  WSA J. Marie Suga  Kaiser Permanente—Baldwin Park  WSA Jonathan A. Polikoff  Kaiser Permanente—Baldwin Park  USA Jonathan A. Polikoff  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  WSA Jonathan A. Polikoff  Marin Cancer Care Inc  WSA Peter D. Eisenberg  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente Medical Center—Vacaville  VSA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale  USA Derrick Wong  University of Colorado Cancer Center  USA Alexander T. Urquhart  Yale University  Smilow Cancer Hospital Care Center—Trumbull  SMCCarron   |   | Seaward  |
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| Kaiser Permanente—Hawaii  Kaiser Permanente—Colorado  Kaiser Permanente—Santa Teresa-San Jose  Saint Joseph's Medical Center  Kaiser Permanente—Santa Teresa-San Jose  Saint Joseph's Medical Center  WSA J. Marie Suga  Saint Joseph's Medical Center  Kaiser Permanente Los Angeles Medical Center  WSA Jonathan A. Polikoff  Kaiser Permanente—Fresno  Sutter Cancer Research Consortium—Sacramento  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Woodland Hills  WSA Jonathan A. Polikoff  Kaiser Permanente—Baldwin Park  WSA Jonathan A. Polikoff  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  WSA Jonathan A. Polikoff  Marin Cancer Care Inc  WSA Jonathan A. Polikoff  Marin Cancer Care Inc  WSA Peter D. Eisenberg  Kaiser Permanente Medical Center—Vacaville  WSA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale  USA Derrick Wong  University of Colorado Cancer Center  WSA Alexander T. Urquhart  Yale University  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospital-Waterbury Care Center  Wedstar Franklin Square Medical Center/Weinberg  USA Erin W. Hofstatter  WedStar Franklin Square Medical Center/Weinberg  USA Edward C. McCarron  | Kaiser Permanente—Southern California   | USA Lara N. Durna  |
| Kaiser Permanente—Colorado Kaiser Permanente—Santa Teresa-San Jose Saint Joseph's Medical Center USA Ajithkumar Puthillath Kaiser Permanente Los Angeles Medical Center USA J. Marie Suga Saint Joseph's Medical Center USA Ajithkumar Puthillath Kaiser Permanente—Fresno USA J. Marie Suga Sutter Cancer Research Consortium—Sacramento USA Nitin Rohatgi Kaiser Permanente—Santa Rosa USA J. Marie Suga USA J. Marie Suga USA J. Marie Suga USA Jonathan A. Polikoff Kaiser Permanente—Woodland Hills USA Jonathan A. Polikoff Contra Costa Regional Medical Center USA James H. Feusner Sutter Cancer Research Consortium—Roseville Kaiser Permanente West Los Angeles USA Jonathan A. Polikoff Marin Cancer Care Inc USA Peter D. Eisenberg Kaiser Permanente Medical Center—Vacaville USA J. Marie Suga Kaiser Permanente Medical Center—Vacaville USA J. Marie Suga USA Jonathan A. Polikoff Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong University of Colorado Cancer Center USA Virginia F. Borges Shaw Cancer Center USA Perin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull USA Erin W. Hofstatter Medstar Franklin Square Medical Center/Weinberg USA Edward C. McCarron   | Kaiser Permanente—Hawaii  |  |
| Kaiser Permanente—Santa Teresa-San Jose Saint Joseph's Medical Center USA Ajithkumar Puthillath  Kaiser Permanente Los Angeles Medical Center USA Jonathan A. Polikoff  Kaiser Permanente—Fresno Sutter Cancer Research Consortium—Sacramento USA J. Marie Suga  Sutter Cancer Research Consortium—Sacramento USA J. Marie Suga  Kaiser Permanente—Santa Rosa USA J. Marie Suga  Kaiser Permanente—Woodland Hills USA Jonathan A. Polikoff  Kaiser Permanente—Baldwin Park USA Jonathan A. Polikoff  Contra Costa Regional Medical Center USA James H. Feusner  Sutter Cancer Research Consortium—Roseville USA Kristie A Bobolis  Kaiser Permanente West Los Angeles USA Jonathan A. Polikoff  Marin Cancer Care Inc USA Peter D. Eisenberg  Kaiser Permanente Medical Center—Vacaville USA J. Marie Suga  Kaiser Permanente—San Marcos USA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong University of Colorado Cancer Center USA Virginia F. Borges  Shaw Cancer Center USA Alexander T. Urquhart  Yale University USA Erin W. Hofstatter  Smilow Cancer Hospital Care Center—Trumbull USA Erin W. Hofstatter  Medstar Franklin Square Medical Center/Weinberg USA Edward C. McCarron   | Kaiser Permanente—Colorado  |  |
| Saint Joseph's Medical Center  Kaiser Permanente Los Angeles Medical Center  Kaiser Permanente—Fresno  Sutter Cancer Research Consortium—Sacramento  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Santa Rosa  Kaiser Permanente—Woodland Hills  Kaiser Permanente—Woodland Hills  Kaiser Permanente—Baldwin Park  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  WSA Jonathan A.  Polikoff  USA Jonathan A.  Polikoff  USA Jonathan A.  Polikoff  Marin Cancer Care Inc  WSA Jonathan A.  Polikoff  Marin Cancer Care Inc  WSA Peter D.  Eisenberg  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente Medical Center—Vacaville  WSA Jonathan A.  Polikoff  Palo Alto Medical Foundation—Sunnyvale  USA Jonathan A.  Polikoff  Palo Alto Medical Foundation—Sunnyvale  USA Derrick Wong  University of Colorado Cancer Center  USA Alexander T.  Urquhart  Yale University  USA Erin W. Hofstatter  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospital-Waterbury Care Center  Wedstar Franklin Square Medical Center/Weinberg  USA Edward C.  McCarron  |   |  |
| Kaiser Permanente Los Angeles Medical Center  Kaiser Permanente—Fresno Sutter Cancer Research Consortium—Sacramento Kaiser Permanente—Santa Rosa USA J. Marie Suga USA Jonathan A. Polikoff  Kaiser Permanente—Baldwin Park USA Jonathan A. Polikoff  Contra Costa Regional Medical Center USA James H. Feusner Sutter Cancer Research Consortium—Roseville USA Kristie A Bobolis Kaiser Permanente West Los Angeles USA Jonathan A. Polikoff  Marin Cancer Care Inc USA Peter D. Eisenberg Kaiser Permanente Medical Center—Vacaville USA J. Marie Suga Kaiser Permanente—San Marcos USA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong University of Colorado Cancer Center USA Virginia F. Borges Shaw Cancer Center USA Alexander T. Urquhart Yale University USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull USA Erin W. Hofstatter Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute  |   | USA Ajithkumar   |
| Sutter Cancer Research Consortium—Sacramento Kaiser Permanente—Santa Rosa Kaiser Permanente—Woodland Hills USA J. Marie Suga Kaiser Permanente—Woodland Hills USA Jonathan A. Polikoff Kaiser Permanente—Baldwin Park USA James H. Feusner Sutter Cancer Research Consortium—Roseville Kaiser Permanente West Los Angeles USA James H. Feusner Sutter Cancer Research Consortium—Roseville Kaiser Permanente West Los Angeles USA Jonathan A. Polikoff Marin Cancer Care Inc USA Peter D. Eisenberg Kaiser Permanente Medical Center—Vacaville USA J. Marie Suga Kaiser Permanente—San Marcos USA Jonathan A. Polikoff Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong University of Colorado Cancer Center USA Virginia F. Borges Shaw Cancer Center USA Alexander T. Urquhart Yale University USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull Smilow Cancer Hospital-Waterbury Care Center USA Erin W. Hofstatter USA Erin W. Hofstatter Medstar Franklin Square Medical Center/Weinberg USA Edward C. McCarron   | Kaiser Permanente Los Angeles Medical Cent  | er USA Jonathan A.   |
| Kaiser Permanente—Santa Rosa Kaiser Permanente—Woodland Hills USA Jonathan A. Polikoff  Kaiser Permanente—Baldwin Park USA Jonathan A. Polikoff  Contra Costa Regional Medical Center Sutter Cancer Research Consortium—Roseville Kaiser Permanente West Los Angeles USA Kristie A Bobolis Kaiser Permanente West Los Angeles USA Jonathan A. Polikoff  Marin Cancer Care Inc USA Peter D. Eisenberg Kaiser Permanente Medical Center—Vacaville Kaiser Permanente—San Marcos USA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong University of Colorado Cancer Center USA Virginia F. Borges Shaw Cancer Center USA Alexander T. Urquhart Yale University USA Erin W. Hofstatter Smilow Cancer Hospital-Waterbury Care Center Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute   | Kaiser Permanente—Fresno  | USA J. Marie Suga  |
| Kaiser Permanente—Woodland Hills  Kaiser Permanente—Baldwin Park  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  Kaiser Permanente West Los Angeles  Was Jonathan A. Polikoff  Marin Cancer Care Inc  USA Peter D. Eisenberg  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente—San Marcos  USA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale  USA Derrick Wong  University of Colorado Cancer Center  USA Alexander T. Urquhart  Yale University  USA Erin W. Hofstatter  Smilow Cancer Hospital-Waterbury Care Center  Medstar Franklin Square Medical Center/Weinberg  USA Edward C. Cancer Institute  | Sutter Cancer Research Consortium—Sacrame   | ento USA Nitin Rohatgi   |
| Raiser Permanente—Baldwin Park  Kaiser Permanente—Baldwin Park  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  Kaiser Permanente West Los Angeles  Marin Cancer Care Inc  Sutter Cancer Care Inc  Was Peter D.  Eisenberg  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente—San Marcos  Was Jonathan A.  Polikoff  Palo Alto Medical Foundation—Sunnyvale  UsA Derrick Wong  University of Colorado Cancer Center  UsA Virginia F. Borges  Shaw Cancer Center  UsA Alexander T.  Urquhart  Yale University  UsA Erin W. Hofstatter  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospital-Waterbury Care Center  Medstar Franklin Square Medical Center/Weinberg  USA Edward C.  McCarron   | Kaiser Permanente—Santa Rosa  | USA J. Marie Suga  |
| Polikoff  Contra Costa Regional Medical Center  Sutter Cancer Research Consortium—Roseville  Kaiser Permanente West Los Angeles  Marin Cancer Care Inc  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente Medical Center—Vacaville  Kaiser Permanente—San Marcos  Kaiser Permanente—San Marcos  USA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale  USA Derrick Wong  University of Colorado Cancer Center  USA Virginia F. Borges  Shaw Cancer Center  USA Alexander T. Urquhart  Yale University  USA Erin W. Hofstatter  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospital-Waterbury Care Center  Medstar Franklin Square Medical Center/Weinberg  USA Edward C. Cancer Institute   | Kaiser Permanente—Woodland Hills  |  |
| Sutter Cancer Research Consortium—Roseville Kaiser Permanente West Los Angeles USA Jonathan A. Polikoff  Marin Cancer Care Inc USA Peter D. Eisenberg Kaiser Permanente Medical Center—Vacaville USA J. Marie Suga USA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong University of Colorado Cancer Center USA Virginia F. Borges Shaw Cancer Center USA Alexander T. Urquhart Yale University USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull Smilow Cancer Hospital-Waterbury Care Center Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute   | Kaiser Permanente—Baldwin Park  |  |
| Kaiser Permanente West Los Angeles  Marin Cancer Care Inc  Marin Cancer Care Inc  WSA Peter D. Eisenberg  Kaiser Permanente Medical Center—Vacaville  WSA J. Marie Suga  Kaiser Permanente—San Marcos  WSA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale  University of Colorado Cancer Center  WSA Virginia F. Borges  Shaw Cancer Center  WSA Alexander T. Urquhart  Yale University  WSA Erin W. Hofstatter  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospital-Waterbury Care Center  Medstar Franklin Square Medical Center/Weinberg  Cancer Institute  USA Edward C. McCarron  | Contra Costa Regional Medical Center  | USA James H. Feusner   |
| Polikoff  Marin Cancer Care Inc  Marin Cancer Care Inc  WSA Peter D. Eisenberg  Kaiser Permanente Medical Center—Vacaville  WSA J. Marie Suga  Kaiser Permanente—San Marcos  USA Jonathan A. Polikoff  Palo Alto Medical Foundation—Sunnyvale  USA Derrick Wong  University of Colorado Cancer Center  USA Virginia F. Borges  Shaw Cancer Center  USA Alexander T. Urquhart  Yale University  USA Erin W. Hofstatter  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospital-Waterbury Care Center  Medstar Franklin Square Medical Center/Weinberg  USA Edward C. Cancer Institute  | Sutter Cancer Research Consortium—Rosevill  | e USA Kristie A Bobolis  |
| Kaiser Permanente Medical Center—Vacaville Kaiser Permanente—San Marcos USA J. Marie Suga Kaiser Permanente—San Marcos USA Jonathan A. Polikoff Palo Alto Medical Foundation—Sunnyvale UsA Derrick Wong University of Colorado Cancer Center USA Virginia F. Borges Shaw Cancer Center USA Alexander T. Urquhart Yale University USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull Smilow Cancer Hospital-Waterbury Care Center Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute  | Kaiser Permanente West Los Angeles  |  |
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| Kaiser Permanente—San Marcos  Polikoff  Palo Alto Medical Foundation—Sunnyvale USA Derrick Wong University of Colorado Cancer Center USA Virginia F. Borges Shaw Cancer Center USA Alexander T. Urquhart  Yale University USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull Smilow Cancer Hospital-Waterbury Care Center Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute  USA Edward C. McCarron   | Kaiser Permanente Medical Center—Vacaville  | •  |
| Palo Alto Medical Foundation—Sunnyvale University of Colorado Cancer Center USA Virginia F. Borges Shaw Cancer Center USA Alexander T. Urquhart Yale University USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull Smilow Cancer Hospital-Waterbury Care Center Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute USA Edward C. McCarron  | Kaiser Permanente—San Marcos  | USA Jonathan A.  |
| University of Colorado Cancer Center  Shaw Cancer Center  USA Virginia F. Borges USA Alexander T. Urquhart  Yale University  USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull Smilow Cancer Hospital-Waterbury Care Center Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute  USA Edward C. McCarron  | Palo Alto Medical Foundation—Sunnyvale  |  |
| Shaw Cancer Center  Yale University  Smilow Cancer Hospital Care Center—Trumbull Smilow Cancer Hospital-Waterbury Care Center  Medstar Franklin Square Medical Center/Weinberg  Cancer Institute  USA Erin W. Hofstatter USA Erin W. Hofstatter USA Erin W. Hofstatter Medstar Franklin Square Medical Center/Weinberg  USA Edward C. McCarron   | ,   |  |
| Yale University  Yale University  Smilow Cancer Hospital Care Center—Trumbull  Smilow Cancer Hospital-Waterbury Care Center  Medstar Franklin Square Medical Center/Weinberg  USA Erin W. Hofstatter  Wedstar Franklin Square Medical Center/Weinberg  USA Edward C.  Cancer Institute   | ,   |  |
| Yale University USA Erin W. Hofstatter Smilow Cancer Hospital Care Center—Trumbull USA Erin W. Hofstatter USA Edward C. Cancer Institute USA Erin W. Hofstatter   |   |  |
| Smilow Cancer Hospital Care Center—Trumbull USA Erin W. Hofstatter Smilow Cancer Hospital-Waterbury Care Center USA Erin W. Hofstatter Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute McCarron   | Vala Hairandita   |  |
| Smilow Cancer Hospital-Waterbury Care Center USA Erin W. Hofstatter Medstar Franklin Square Medical Center/Weinberg USA Edward C. Cancer Institute McCarron  | Yale University   | USA Erin W. Hofstatter   |
| Medstar Franklin Square Medical Center/Weinberg USA Edward C.<br>Cancer Institute McCarron   | ,   |  |
| Cancer Institute McCarron  | Smilow Cancer Hospital Care Center—Trumbu   | ull USA Erin W. Hofstatter   |
|  | Smilow Cancer Hospital Care Center—Trumbo<br>Smilow Cancer Hospital-Waterbury Care Cent   | ull USA Erin W. Hofstatter<br>er USA Erin W. Hofstatter                                      |
|  | Smilow Cancer Hospital Care Center—Trumbu<br>Smilow Cancer Hospital-Waterbury Care Cent<br>Medstar Franklin Square Medical Center/Wei                     | ull USA Erin W. Hofstatter<br>ter USA Erin W. Hofstatter<br>nberg USA Edward C.              |
| Continued  | Smilow Cancer Hospital Care Center—Trumbu<br>Smilow Cancer Hospital-Waterbury Care Cent<br>Medstar Franklin Square Medical Center/Wei<br>Cancer Institute | ull USA Erin W. Hofstatter<br>ter USA Erin W. Hofstatter<br>inberg USA Edward C.<br>McCarron |

| Appendix Continued  |     |                                      |
|---|-----|--------------------------------------|
| MedStar Washington Hospital Center  | USA | Pia<br>Herbolsheimer                 |
| ChristianaCare Oncology Hematology at the Helen   | USA |                                      |
| F. Graham Cancer Center & Research Institute  |     | Varadarajan                          |
| Helen F Graham Cancer Center & Research Institute   |     |                                      |
| Halifax Health Medical Center-Centers for Oncology  | USA | Deveras                              |
| University of Miami Miller School of<br>Medicine—Sylvester Cancer Center                            |     | Frances Valdes-<br>Albini            |
| UM Sylvester Comprehensive Cancer Center at Deerfield Beach   |     | Reshma L.<br>Mahtani                 |
| UM Sylvester Comprehensive Cancer Center at Plantation  |     | Reshma L.<br>Mahtani                 |
| Emory University Hospital/Winship Cancer Institute<br>Medical Center of Central Georgia             |     | Jane L. Meisel<br>Bradley T. Sumrall |
| Northside Hospital, Georgia NCORP   |     | Cheryl F. Jones                      |
| South Georgia Medical Center/Pearlman Cancer  |     | Samuel N. Ofori                      |
| Center<br>Straub Clinic and Hospital  | USA | Kenneth N.M.                         |
| Pali Momi Medical Center  | USA | Sumida<br>Kenneth N.M.               |
|   |     | Sumida<br>Mark Karwal                |
| University of Iowa/Holden Comprehensive Cancer<br>Center  |     |                                      |
| Oncology Associates at Mercy Medical Center   | USA | Deborah W.<br>Wilbur                 |
| Mercy Medical Center—North Iowa   | USA | Joginder (Joe)<br>Singh              |
| Genesis Medical Center—East Campus  | USA | David M. Spector                     |
| Kootenai Cancer Center  | USA | John<br>Schallenkamp                 |
| NorthShore University HealthSystem-Highland Park Hospital   | USA | Douglas E.<br>Merkel                 |
| Loyola University Medical Center  | USA | Shelly S. Lo                         |
| Mount Sinai Hospital Medical Center   |     | Pam G. Khosla                        |
| Northwestern University   | USA | Massimo<br>Cristofanilli             |
| Northwestern Medicine, Robert H. Lurie<br>Comprehensive Cancer Center of Northwestern<br>University | USA | Lisa Flaum                           |
| University of Illinois  | USA | Kent F. Hoskins                      |
| Rush University Medical Center  |     | Melody A.<br>Cobleigh                |
| Swedish Covenant Hospital   | USA | Elyse A. Lambiase                    |
| University of Chicago Comprehensive Cancer Center   |     |                                      |
| Presence Saint Joseph Hospital—Chicago  |     | Ira A. Oliff                         |
| Missouri Baptist Cancer Center  |     | Bryan A. Faller                      |
| Illinois CancerCare—Peoria  |     | James L. Wade                        |
| Joliet Oncology-Hematology Associates Limited   |     | Nafisa D. Burhan                     |
| Cancer Care Specialists of Illinois—Decatur Elmhurst Memorial Hospital                              |     | James L. Wade<br>Amaryllis Gil       |
| SwedishAmerican Regional Cancer Center  |     | Harvey E. Einhorn                    |
| Indiana University School of Medicine/Melvin and  |     | Anna M.V.                            |
| Bren Simon Cancer Center  |     | Storniolo                            |
| Parkview Hospital Randallia   |     | Brian K. Chang                       |
| IU Health Ball Memorial Hospital  |     | Maitri Kalra                         |
| The Community Hospital Michiana Hematology Oncology PC—Mishawaka                                    |     | Erwin L. Robin<br>Bilal Ansari       |
| Department of Internal Medicine, Division Medical   |     | Priyanka Sharma                      |
| Oncology, University of Kansas Medical Center<br>Cancer Center of Kansas—Wichita                    |     | Shaker R. Dakhil                     |
| Cancer Center of Kansas-Wichita Medical Arts<br>Tower   |     | Shaker R. Dakhil                     |
| Saint Joseph Hospital East  | USA | Richard L.<br>Deming                 |
| Ochsner Medical Center Jefferson  |     | John T. Cole                         |
| CHRISTUS Highland Medical Center  |     | John T. Cole                         |
| Ochsner Health Center—Summa   |     | John T. Cole                         |
| Our Lady of the Lake Physician Group  |     | David S. Hanson                      |
| Louisiana Hematology Oncology Associates LLC  |     | Augusto C. Ochoa                     |
| Ochsner Medical Center Kenner  Many Rind Parkins Capper Center Covington                            |     | John T. Cole                         |
| Mary Bird Perkins Cancer Center—Covington   | USA | Augusto C. Ochoa                     |
|   |     | Continued                            |

| Appendix Continued  |     |                                    |
|---|-----|------------------------------------|
| Dana-Farber/Harvard Cancer Center   | USA | Judy E. Garber                     |
| Beth Israel Deaconess Medical Center  | USA | Judy E. Garber                     |
| Berkshire Medical Center—Cancer Center  | USA | Harvey Zimbler                     |
| Suburban Hospital   | USA | Deborah K.<br>Armstrong            |
| University of Maryland/Greenebaum Cancer Center                                       | USA | •                                  |
| Mercy Medical Center  | USA | David A. Riseberg                  |
| Johns Hopkins University/Sidney Kimmel Cancer   | USA | Deborah K.                         |
| Center  |     | Armstrong                          |
| Frederick Memorial Hospital   | USA | Brian M.<br>O'Connor               |
| Eastern Maine Medical Center  | USA | Thomas H.<br>Openshaw              |
| Penobscot Bay Medical Center  | USA | Thomas H.<br>Openshaw              |
| Harold Alfond Center for Cancer Care  | USA | Thomas H.<br>Openshaw              |
| William Beaumont Hospital—Royal Oak   | USA | Dana Zakalik                       |
| Ascension Providence Hospitals—Southfield   | USA | Cynthia M.                         |
|   |     | Vakhariya                          |
| University of Michigan Rogel Cancer Center  |     | Anne F. Schott                     |
| Wayne State University/Karmanos Cancer Institute                                      |     | Michael S. Simon                   |
| Henry Ford Hospital   |     | Thomas J. Doyle                    |
| Trinity Health Ann Arbor Hospital, Michigan Cancer                                    | USA |                                    |
| Research Consortium (NCORP)   |     | Baghdadi                           |
| Cancer Research Consortium of West Michigan,<br>Spectrum Health at Butterworth Campus | USA | Amy<br>VanderWoude                 |
| Regions Hospital  | USA | Patrick J. Flynn                   |
| Mercy Hospital  | USA | Richard T. Zera                    |
| Essentia Health Cancer Center   | USA | Bret E.B. Friday                   |
| Mayo Clinic   | USA | Kathryn J. Ruddy                   |
| Saint Francis Regional Medical Center   | USA | Richard T. Zera                    |
| Mayo Clinic Health Systems—Mankato  |     | Ron Smith                          |
| Fairview Clinics and Surgery Center Maple Grove                                       |     | Patrick J. Flynn                   |
| Washington University School of Medicine  | USA | Foluso Olabisi<br>Ademuyiwa        |
| CoxHealth South Hospital  |     | Robert Ellis                       |
| Mercy Hospital Springfield  |     | Jay W. Carlson                     |
| Saint Louis Cancer and Breast Institute—South City                                    |     |                                    |
| Kalispell Regional Medical Center   |     | Marchello,<br>Benjamin T.          |
| Atrium—Wake Forest University Health Sciences Duke University Medical Center          |     | Edward A. Levine<br>Paul K. Marcom |
| Mission Hospital  |     | Cameron B.                         |
| IVIISSIOII HOSPITAI   | USA | Harkness                           |
| Levine Cancer Institute, Atrium Health  | USA | Antoinette R. Tan                  |
| CaroMont Regional Medical Center  |     | William J. Charles                 |
| FirstHealth of the Carolinas—Pinehurst  |     | Charles S. Kuzma                   |
| Southeastern Medical Oncology   |     | Shonda Asaad                       |
| Center—Jacksonville   |     |                                    |
| Margaret R Pardee Memorial Hospital   |     | James E. Radford                   |
| Sanford Roger Maris Cancer Center   |     | Preston D. Steen                   |
| Trinity Cancer Care Center  | USA | Madhu<br>Unnikrishnan              |
| Altru Cancer Center   | USA | Grant R. Seeger                    |
| Nebraska Methodist Hospital   | USA | Kirsten M.H. Leu                   |
| CHI Health Saint Francis  | USA | Mehmet S. Copur                    |
| Southeast Nebraska Cancer Center—68th Street Place                                    | USA | Ralph J. Hauke                     |
| Nebraska Hematology and Oncology  | USA | Gamini S. Soori                    |
| Faith Regional Health Services Carson Cancer Center                                   | USA | Ralph J. Hauke                     |
| Dartmouth-Hitchcock Medical Center/Norris Cotton                                      | USA | Bradley A. Arrick                  |
| Cancer Center Morristown Medical Center   | USA | Jennifer G.                        |
| Rutgers Cancer Institute of New Jersey  | USA | Reeder<br>Deborah L.               |
|   |     | Toppmeyer                          |
| University of New Mexico Comprehensive Cancer<br>Center (NM MU-NCORP)                 |     | Zoneddy R.<br>Dayao                |
| Laura and Isaac Perlmutter Cancer Center at NYU<br>Langone                            | USA | Sylvia Adams                       |
|   |     | Continued                          |

| NYP/Weill Cornell Medical Center   | USA                                    | Eleni<br>Andreopoulou   |
|--|--|---|
| University of Rochester  | USA                                    | Magnuson<br>Allison   |
| Montefiore Medical Center—Einstein Campus  |  | Jesus D. Anampa<br>Mesias   |
| Northwell Health Cancer Institute Ohio State University Comprehensive Cancer Center  |  | Ruby Sharma<br>Bhuvaneswari<br>Ramaswamy  |
| Cleveland Clinic Foundation  | USA                                    | Aaron T. Gerds  |
| UH Seidman Cancer Center at Southwest General Hospital   |  | Robert R. Shenk   |
| Kettering Medical Center Aultman Health Foundation   |  | Howard M. Gross<br>Shruti Trehan  |
| Miami Valley Hospital North  | USA                                    | Howard M. Gross   |
| Blanchard Valley Hospital  |  | Howard M. Gross   |
| Dayton Physician LLC-Miami Valley Hospital North   |  | Howard M. Gross   |
| UHHS-Chagrin Highlands Medical Center  |  | Robert R. Shenk   |
| Springfield Regional Cancer Center Mercy Cancer Center-Elyria  |  | Howard M. Gross<br>Robert R. Shenk  |
| Stephenson Cancer Center, University of Oklahoma<br>Health Sciences Center   |  |   |
| Kaiser Permanente Northwest  | USA                                    | Abdul H.<br>Mansoor   |
| Allegheny Health Network   | USΔ                                    | Christie J. Hilton  |
| UPMC Hillman Cancer Center   |  | Adam M. Brufsky   |
| WellSpan Health  |  | Chanh Huynh   |
| Delaware County Memorial Hospital  | USA                                    | Nabila<br>Chowdhury   |
| Basser Center for BRCA at the Abramson Cancer<br>Center, University of Pennsylvania  | USA                                    | Susan M.<br>Domchek   |
| Fox Chase Cancer Center  | USA                                    | Elin R. Sigurdson   |
| Reading Hospital   | USA                                    | Terrence P.<br>Cescon   |
| Penn State Health Saint Joseph Medical Center  |  | Marc A. Rovito  |
| Lankenau Medical Center  |  | Albert S. DeNittis  |
| Geisinger Wyoming Valley/Henry Cancer Center   |  | Victor G. Vogel   |
| Jefferson Hospital Adams Cancer Center   |  | Thomas B. Julian<br>L. E. Boyle   |
| San Juan City Hospital   |  | Luis Baez-Diaz  |
| Medical University of South Carolina   |  | Frank J. Brescia  |
| AnMed Health Cancer Center   | USA                                    | John E. Doster  |
| Saint Francis Cancer Center  |  | Robert D. Siegel  |
| Scott and White Memorial Hospital  |  | Lucas Wong  |
| Houston Methodist Hospital   |  | Tejal Patel   |
| Baylor College of Medicine/Dan L Duncan<br>Comprehensive Cancer Center   |  | Julie R. Nangia   |
| Texas Tech University Health Sciences Center—Lubbock   |  | Catherine A. Jones  |
| McKay-Dee Hospital Center  |  | George M.<br>Cannon   |
| Utah Valley Regional Medical Center  |  | George M.<br>Cannon   |
| Virginia Commonwealth University/Massey Cancer   |  | Harry D. Bear   |
| Center   |  |   |
| Virginia Commonwealth University/Massey Cancer Center  |  | Hetal Vachhani  |
| Virginia Commonwealth University/Massey Cancer<br>Center<br>Inova Schar Cancer Institute   | USA                                    | Mary Wilkinson  |
| Virginia Commonwealth University/Massey Cancer<br>Center<br>Inova Schar Cancer Institute<br>University of Vermont and State Agricultural College   | USA<br>USA                             | Mary Wilkinson<br>Marie E. Wood   |
| Virginia Commonwealth University/Massey Cancer<br>Center<br>Inova Schar Cancer Institute   | USA<br>USA<br>USA                      | Mary Wilkinson<br>Marie E. Wood<br>Marie E. Wood  |
| Virginia Commonwealth University/Massey Cancer<br>Center<br>Inova Schar Cancer Institute<br>University of Vermont and State Agricultural College<br>Central Vermont Medical Center<br>Swedish Medical Center—First Hill  | USA<br>USA<br>USA<br>USA               | Mary Wilkinson<br>Marie E. Wood<br>Marie E. Wood<br>Fengting Yan  |
| Virginia Commonwealth University/Massey Cancer<br>Center<br>Inova Schar Cancer Institute<br>University of Vermont and State Agricultural College<br>Central Vermont Medical Center<br>Swedish Medical Center—First Hill<br>Providence Regional Cancer System—Centralia<br>University of Washington School of Medicine,<br>Division of Oncology Fred Hutch/University of  | USA<br>USA<br>USA<br>USA<br>USA        | Mary Wilkinson<br>Marie E. Wood<br>Marie E. Wood  |
| Virginia Commonwealth University/Massey Cancer Center Inova Schar Cancer Institute University of Vermont and State Agricultural College Central Vermont Medical Center Swedish Medical Center—First Hill Providence Regional Cancer System—Centralia University of Washington School of Medicine, Division of Oncology Fred Hutch/University of Washington Cancer Consortium University of Washington School of Medicine, Division of Oncology Fred Hutch/University of  | USA<br>USA<br>USA<br>USA<br>USA        | Mary Wilkinson<br>Marie E. Wood<br>Marie E. Wood<br>Fengting Yan<br>Xingwei Sui<br>Carol M. van                                     |
| Virginia Commonwealth University/Massey Cancer Center Inova Schar Cancer Institute University of Vermont and State Agricultural College Central Vermont Medical Center Swedish Medical Center—First Hill Providence Regional Cancer System—Centralia University of Washington School of Medicine, Division of Oncology Fred Hutch/University of Washington Cancer Consortium University of Washington School of Medicine,  | USA<br>USA<br>USA<br>USA<br>USA<br>USA | Mary Wilkinson<br>Marie E. Wood<br>Marie E. Wood<br>Fengting Yan<br>Xingwei Sui<br>Carol M. van<br>Haelst<br>Jennifer M.            |
| Virginia Commonwealth University/Massey Cancer Center Inova Schar Cancer Institute University of Vermont and State Agricultural College Central Vermont Medical Center Swedish Medical Center—First Hill Providence Regional Cancer System—Centralia University of Washington School of Medicine, Division of Oncology Fred Hutch/University of Washington Cancer Consortium University of Washington School of Medicine, Division of Oncology Fred Hutch/University of Washington Cancer Consortium                                       | USA<br>USA<br>USA<br>USA<br>USA<br>USA | Mary Wilkinson Marie E. Wood Marie E. Wood Fengting Yan Xingwei Sui Carol M. van Haelst Jennifer M. Specht                          |
| Virginia Commonwealth University/Massey Cancer Center Inova Schar Cancer Institute University of Vermont and State Agricultural College Central Vermont Medical Center Swedish Medical Center—First Hill Providence Regional Cancer System—Centralia University of Washington School of Medicine, Division of Oncology Fred Hutch/University of Washington Cancer Consortium University of Washington School of Medicine, Division of Oncology Fred Hutch/University of Washington Cancer Consortium Kadlec Clinic Hematology and Oncology | USA<br>USA<br>USA<br>USA<br>USA<br>USA | Mary Wilkinson Marie E. Wood Marie E. Wood Fengting Yan Xingwei Sui Carol M. van Haelst  Jennifer M. Specht  Ying Zhuo Rubina Qamar |

| Appendix Continued   |                            |
|--|----------------------------|
| Aurora Cancer Care-Southern Lakes VLCC                     | USA Shamsuddin<br>Virani   |
| Aurora BayCare Medical Center                              | USA Rubina Qamar           |
| Marshfield Medical Center—Weston                           | USA Arlene A. Gayle        |
| Aurora Cancer Care—Grafton                                 | USA Rubina Qamar           |
| Aurora Health Center—Fond du Lac                           | USA Rubina Qamar           |
| Charleston Area Medical Center, David Lee Cancer<br>Center | USA Steven J.<br>Jubelirer |
| West Virginia University Cancer Institute                  | USA Sobha Kurian           |
| West Virginia University Healthcare                        | USA Mohamad A.<br>Salkeni  |

## SABO: Swedish Association of Breast Oncologists

| Skånes Universitetssjukhus Lund/Skåne/Lund<br>University Hospital, Department of Oncology,<br>Malmö | Sweden Niklas Loman                        |
|---|--|
| Sahlgrenska Universitetssjukhuset,<br>Gothenburg  | Sweden Barbro Linderholm                   |
| Norrlands Universitetssjukhus, Umeå   | Sweden Gustav Silander                     |
| Linköpings Universitetssjukhus, Linköping   | Sweden Anna-Lotta<br>Hallbeck              |
| Södersjukhuset, Stockholm   | Sweden Anna von<br>Wachenfeldt<br>Väppling |

## SOLTI

| Hôpital Jean Minjoz                                       | France   | Elsa Curtit                    |
|---|----------|--------------------------------|
| IPO Lisboa, Serviço de Oncologia Médica 2                 | Portugal | Catarina Cardoso               |
| Hospital CUF Descobertas                                  | Portugal | Sofia Braga                    |
| IPO Porto, Serviço de Oncologia Médica                    | Portugal | Miguel Abreu                   |
| Hospital Beatriz Ângelo, Hospital de Dia<br>Oncologia     | Portugal | Mafalda Casa-Nova              |
| Hospital da Luz   | Portugal | Mónica Nave                    |
| Hospital Universitario 12 de Octubre                      | Spain    | Eva María Ciruelos Gil         |
| Hospital Vall d'Hebron                                    | Spain    | Judith Balmaña Gelpi           |
| Institut Catala d'Oncologia Hospitalet                    | Spain    | Adela Fernández<br>Ortega      |
| Hospital San Joan de Reus                                 | Spain    | Josep Gumà Padró               |
| Hospital Clínico Universitario de Valencia                | Spain    | Begoña Bermejo de<br>las Heras |
| Usp Institut Universitari Dexeus                          | Spain    | María González Cao             |
| Complejo Hospitalario Universitario de<br>Santiago (CHUS) | Spain    | Juan Cueva Bañuelos            |
| Hospital Universitario Son Espases                        | Spain    | Jesús Alarcon<br>Company       |
| Hospital Josep Trueta                                     | Spain    | Gemma Viñas Villaró            |
| MD Anderson Cancer Center                                 | Spain    | Laura García Estevez           |
|   |          |                                |

## **SUCCESS**

| Universitätsklinikum Ulm                        | Germany Jens Huober               |
|---|-----------------------------------|
| Brustzentrum Mittelthüringen                    | Germany Steffi Busch              |
| Universitätsklinikum Düsseldorf                 | Germany Tanja Fehm                |
| Stadtklinik Baden-Baden                         | Germany Antje Hahn                |
| Südharz-Krankenhaus Nordhausen gGmbH            | Germany Andrea Grafe              |
| Kreiskrankenhaus Hameln                         | Germany Thomas Noesselt           |
| Klinikum Gifhorn GmbH                           | Germany Thomas Dewitz             |
| Gemeinschaftspraxis Drs. med. Wilke/Wagner      | Germany Harald Wagner             |
| Klinikum Memmingen                              | Germany Christina Bechtner        |
| Leopoldina-Krankenhaus der Stadt<br>Schweinfurt | Germany Michael Weigel            |
| Marienhospital Bottrop gGmbH                    | Germany Hans-Christian<br>Kolberg |
| Onkologie Ravensburg                            | Germany Thomas Decker             |
|   | Continued                         |

| Appendix Continued                                      |         |                        |
|---|---------|------------------------|
| Institut für Versorgungsforschung in der Onkologie      | Germany | Jörg Thomalla          |
| Diakoniekrankenhaus Rotenburg (Wümme) gGmbH             | Germany | Tobias Hesse           |
| Klinikum der Ludwig-Maximillians-Universität<br>München | Germany | Nadia Harbeck          |
| Onkologische Schwerpunktpraxis Mülheim                  | Germany | Jan Schröder           |
| Charité—Universitätsmedizin Berlin                      | Germany | Jens-Uwe Blohmer       |
| Universitätsklinikum Mannheim                           | Germany | Marc Wolf<br>Sütterlin |
| SweBCG Swedish Breast Cancer Group                      |         |                        |
| Karolinska Universitetssjukhuset, Solna                 | Sweden  | Renske Altena          |

## TCOG: Taiwan Cooperative Oncology Group

| China Medical University Hospital                          | Taiwan Chang-Fang Chiu      |
|--|-----------------------------|
| Chang-Gung Medical Foundation Linkou                       | Taiwan Shin-Cheh Chen       |
| Kaohsiung Medical University Chung-Ho<br>Memorial Hospital | Taiwan Ming-Feng Hou        |
| Mackay Memorial Hospital                                   | Taiwan Yuan-Ching Chang     |
| Chi Mei Hospital-Liou Yin                                  | Taiwan Shang-Hung Chen      |
| Changhua Christian Hospital                                | Taiwan Shou-Tung Chen       |
| National Taiwan University Hospital                        | Taiwan Chiun-Sheng<br>Huang |
| Veterans General Hospital Taichung                         | Taiwan Dah-Cherng Yeh       |
| Triple Service General Hospital                            | Taiwan Jyh-Cherng Yu        |
| Veteran General Hospital Taipei                            | Taiwan Ling-Ming Tseng      |
| National Cheng Kung University (NCKU)<br>Hosptial          | Taiwan Wei-Pang Chung       |

## **UCBG:** Unicancer Breast Group

| Centre Oscar Lambret                   | France Audrey Mailliez               |
|--|--------------------------------------|
| Centre Paul Strauss                    | France Thierry Petit                 |
| Institut Gustave Roussy                | France Suzette DELALOGE              |
| Centre François Baclesse               | France Christelle Lévy               |
| Hôpital Européen de Marseille          | France Philippe Dalivoust            |
| Institut Paoli Calmettes               | France Jean-Marc Extra               |
| Centre Jean Perrin                     | France Marie-Ange Mouret-<br>Reynier |
| Centre CARIO-HPCA                      | France Anne-Claire Hardy-<br>Bessard |
| CHU Morvan-Institut de Cancerologie et | France Hélène Simon                  |
| d'Hematologie                          |                                      |
| Centre Hospitaliser Départemental Les  | France Tiffenn L'Haridon             |
| Oudairies                              |                                      |
| Institut Sainte Catherine              | France Alice Mege                    |
| Hôpital Saint Louis                    | France Sylvie Giacchetti             |
| Institut Bergonié                      | France Camille Chakiba-              |
|  | Brugere                              |
| Clinique Pasteur                       | France Alain Gratet                  |
| Centre Léonard de Vinci                | France Virginie Pottier              |
| Centre Antoine Lacassagne              | France Jean-Marc FERRERO             |
| Centre Henri Becquerel                 | France Isabelle Tennevet             |
| Centre Eugène Marquis                  | France Christophe Perrin             |

## **Independent Sites**

| •  |         |                   |
|--|---------|-------------------|
| Grand Hôpital de Charleroi (GHdC)                        | Belgium | Jean-Luc<br>Canon |
| Universitair Ziekenhuis Brussel                          | Belgium | Sofie Joris       |
| Fudan University Shanghai Cancer Center                  | China   | Zhimin Shao       |
| Cancer Hospital, CAMS&PUMC                               | China   | Binghe Xu         |
| PLA 307 hospital   | China   | ZeFei Jiang       |
| Peking Union Medical College Hospital                    | China   | Qiang Sun         |
| Ruijin hospital Shanghai Jiaotong University of medicine | China   | Kunwei Shen       |
|  |         | Continued         |

| Appendix Continued   |                      |                               |
|--|----------------------|-------------------------------|
| Harbin Medical University Cancer Hospital  | China                | Da Pang                       |
| Tianjin Medical University Cancer Institute and Hospital   |                      | Jin Zhang                     |
| Jiangsu Province Hospital  | China                | Shui Wang                     |
| The Cancer Hospital of the University of Chinese<br>Academy of Sciences (Zhejiang Cancer Hospital)<br>Institute of Basic Medicine and Cancer (IBMC),<br>Chinese Academy of Sciences. |                      | Hongjian Yang                 |
| Guangdong Provincial People's Hospital   | China                | Ning Liao                     |
| West China Hospital, Sichuan University  | China                | Hong Zheng                    |
| The 1st Affiliated Hospital of Medical School of Zhejiang Un The Union Hospital affiliated to Fujian Medical   | China                | Peifen Fu<br>Chuangui         |
| University   |                      | Song                          |
| ShanDong Cancer Hospital   | China                | Yongsheng<br>Wang             |
| The First Hospital of Jilin University   | China                | Zhimin Fan                    |
| Hebei Medical University Fourth Hospital   | China                | Cuizhi Geng                   |
| Centre Léon Bérard   | France               | Olivier Tredan                |
| Uzsoki utcai Kórház  | Hungary              | László<br>Landherr            |
| Chaim Sheba Medical Centre at Tel Hashomer   | Israel               | Bella<br>Kaufman <sup>a</sup> |
| Institute of Oncology, Davidoff Cancer Center,<br>Rabin Medical Center, Beilinson Hospital, Petach<br>Tikva and Sackler Faculty of Medicine, Tel Aviv<br>University, Tel Aviv        | Israel               | Rinat<br>Yerushalmi           |
| Hadassah Hebrew University Medical Center  | Israel               | Beatrice<br>Uziely            |
| Istituto Oncologico Veneto   | Italy                | Pierfranco<br>Conte           |
| A.O.U. di Bologna—Policlinico Sant'Orsola-<br>Malpighi   | Italy                | Claudio<br>Zamagni            |
| Ospedale S. Raffaele—Milano  | Italy                | Giampaolo<br>Bianchini        |
| Istituto Nazionale Tumori Fondazione Pascale IRCCS   | Italy                | Michelino De<br>Laurentiis    |
| Ospedali Riuniti—Azienda Ospedaliera Papa<br>Giovanni XXIII  | Italy                | Carlo Tondini                 |
| La Maddalena Clinic For Cancer University Of<br>Palermo  | Italy                | Vittorio<br>Gebbia            |
| Azienda Ospedaliera Vito Fazzi   | Italy                | Mariangela<br>Ciccarese       |
| Magodent Szpital Elblaska<br>Î   | Poland               | Tomasz<br>Sarosiek            |
| Med Polonia Sp.Z.o.o NSZOZ   | Poland               | Jacek<br>Mackiewicz           |
| SPZOZ MSWiA z Warmińsko-Mazurskim<br>Centrum Onkologii   | Poland               | Anna<br>Słowińska             |
| Instytut Centrum Zdrowia Matki Polki   | Poland               | Ewa Kalinka                   |
| Niepubliczny Zakład Opieki Zdrowotnej<br>Innowacyjna Medycyna  | Poland               | Tomasz<br>Huzarski            |
| Seoul National University Hospital   | Republic of Korea    | Seock-Ah Im                   |
| Asan Medical Center  | Republic of Korea    | Kyung Hae<br>Jung             |
| Yonsei University Severance Hospital   | Republic<br>of Korea | Joo Hyuk Sohn                 |
| Seoul National University Bundang Hospital   | Republic of Korea    | Jee Hyun Kim                  |
| National Cancer Center   | Republic<br>of Korea | Keun Seok Lee                 |
| Samsung Medical Center   | Republic of Korea    | Yeon Hee Park                 |
| Ewha Womans University Mokdong Hospital  | Republic<br>of Korea | Kyoung Eun<br>Lee             |
| Chilgok Kyungpook National University Medical<br>Center  | Republic of Korea    | Yee Soo Chae                  |
| Gachon University Gil Hospital   | Republic<br>of Korea | Eun Kyung<br>Cho              |
| 3-   |                      |                               |

<sup>a</sup>Deceased.