Factors associated with enrolment in clinical trials among women with early-stage breast cancer

D. Presti1, J. Havas1, D. Soldato1,2, P. Lapidari1,3, E. Martin1, B. Pistilli4, C. Jouannaud1, G. Emile6, O. Rigal7, M. Fournier8, P. Soulie9, M.-A. Mouret-Reynier10, C. Tarpin11, M. Campone12, S. Guillermet13, A.-L. Martin14, S. Everhard14 & A. Di Meglio1*

1INSERM Unit 981 - Molecular Predictors and New Targets in Oncology, Gustave Roussy, Villejuif, France; 2Department of Internal Medicine and Medical Specialties (DiMI), School of Medicine, University of Genova, Genoa; 3Department of Internal Medicine and Therapeutics, University of Pavia, Pavia, Italy; 4Department of Medical Oncology, Gustave Roussy, Villejuif; 5Institut Jean Godinot, Reims; 6Centre François Baclesse, Caen; 7Centre Henri Becquerel, Rouen; 8Institut Bergonié, Bordeaux; 9Institut de Cancérologie de l'Ouest - Paul Papin, Angers; 10Centre Jean-Perrin, Clermont Ferrand; 11Institut Paoli Calmettes, Marseille; 12Institut de Cancérologie de l'Ouest - Site de Nantes - Centre René Gauducheau, Nantes; 13Centre Eugene Marquis, Rennes; 14UNICANCER, Paris, France

Available online 17 June 2022

Background: Clinical trials allow development of innovative treatments and ameliorate the quality of clinical care in oncology. Data show that only a minority of patients are enrolled in clinical trials. We assessed enrolment in clinical trials and its correlates among women with early breast cancer.

Methods: We included 9516 patients with stage I-III breast cancer from the multicenter, prospective CANTO study (NCT01993498), followed-up until year 4 (Y4) post-diagnosis. We assessed factors associated with enrolment using multivariable logistic regression. In exploratory, propensity score matched analyses, we used multiple linear regression to evaluate the relationship of enrolment in clinical trials with the European Organisation for Research and Treatment of Cancer Quality Of Life (QoL) questionnaire (EORTC QLQ-C30) Summary Score and described clinical outcomes (distant disease event, invasive disease event, and death by any cause) according to enrolment.

Results: Overall, 1716 patients (18%) were enrolled in a clinical trial until Y4 post-diagnosis of breast cancer. Socioeconomic factors were not associated with enrolment. Centres of intermediate volume were most likely to enrol patients in clinical trials [versus low volume, odds ratio 1.45 (95% confidence interval (CI) 1.08-1.95), \( P = 0.0124 \)]. Among 2118 propensity score matched patients, enrolment was associated with better QoL at Y4 (adjusted mean difference versus not enrolled 1.37, 95% CI 0.03-2.71, \( P = 0.0458 \)), and clinical outcomes (enrolled versus not enrolled, distant disease event 7.3% versus 10.1%, \( P = 0.0206 \); invasive disease event 8.2% versus 10.5%, \( P = 0.0732 \); death by any cause 2.8% versus 3.7%, \( P = 0.2707 \)).

Conclusions: In this large study, one in five patients enrolled on a clinical trial until Y4 after diagnosis of early breast cancer. Geographical and centre-related factors were significantly associated with enrolment in clinical trials. Inclusion in clinical trials seemed associated with improved QoL and clinical outcomes. Access to innovation for early-stage breast cancer patients should be encouraged and facilitated by overcoming organizational and geographical barriers to recruitment.

Key words: breast neoplasms, clinical trial, survivorship, quality of life

INTRODUCTION

Clinical trials are the backbone and foundation of modern evidence-based oncology, as they represent a fundamental instrument to develop innovative treatments and test strategies that can optimize the quality of clinical care. Literature suggests how being enrolled in clinical trials may improve survival and reduce morbidity in patients with cancer of all types, thanks to early access to the latest and most promising investigational interventions, and close clinical monitoring.1-4 Participation in clinical trials and equal access to innovation represent, therefore, key aspects in modern oncology care. According to the National Comprehensive Cancer Network, ‘the best management for any patient with cancer is in a clinical trial’.5 Nevertheless, it has been estimated that only 3%-5% of eligible adult cancer patients in the United States6 and 2%-11% in Australia7 are
enrolled in clinical trials, mostly at the time of cancer recurrence. Trends regarding clinical trial enrolment seem not to have changed substantially over time, suggesting that barriers to trial participation are still numerous and sometimes difficult to overcome, and that access to innovation may be still suboptimal.\(^6,8\) Several studies tried to elucidate the reasons for low recruitment in clinical trials, usually giving emphasis to patient-related barriers, and evaluating patients’ attitude and motivations about enrolment.\(^6,10\) Fear of delay of initiation of antineoplastic drugs, or fear of being enrolled in the placebo arm due to randomization procedures were identified as major drivers of non-participation.\(^11\) In addition, \(\sim 85\%\) of patients receive treatment in community or in small-volume facilities of care, whereas only \(\sim 15\%\) are treated in larger, urban academic centres, where the number of available protocols is generally higher.\(^12\) Socioeconomic and demographic issues can also influence patients’ decision making regarding clinical trial participation. For example, participation in a clinical trial usually requires an increased frequency of follow-up visits, more testing, and more procedures, possibly leading to concerns about travel costs, particularly among patients that have longer travel times to their facility of care or lower income.\(^13\) On the contrary, some factors were found to be associated with higher rates of enrolment in clinical trials, including younger age, male sex, Caucasian race, and having later-stage cancer.\(^14\) Although some factors that may act as barriers or facilitators to clinical trial enrolment were identified, limitations of prior studies include the lack of an extensive characterization of clinical, socioeconomic, tumour, and treatment-related information that may represent relevant determinants of enrolment. Furthermore, previous studies were often single-institution experiences.\(^2,4,15,16\) In addition, evidence regarding determinants of enrolment in clinical trials among patients with early-stage breast cancer is still limited.

In the present analysis, we specifically aimed to examine factors associated with enrolment in clinical trials among women with early breast cancer, using data from the CANcer TOxicity cohort (CANTO). In addition, we assessed the relationship of enrolment in clinical trials with long-term (i) patient-reported functional health and symptom burden, and (ii) survival outcomes.

**PATIENTS AND METHODS**

**Data source**

CANTO (ClinicalTrials.gov/NCT01993498) is a prospective observational cohort study that includes patients with stage I-III breast cancer across 26 participating French cancer centres since 2012. CANTO recruiting centres include 20 comprehensive cancer centres, two university hospitals in Paris, two public non-teaching hospitals and two private hospitals (Supplementary Figure S1, available at https://doi.org/10.1016/j.esmoop.2022.100513). UNICANCER, the National Cooperative Group of French Cancer Centers, coordinates CANTO. The CANTO study was approved by the national regulatory authorities on 14 September 2011 (ref 2011-A011095-36) and the central ethical committee for human subjects (CPP - Ile de France 7, on 14 October 2011—ref 11-039). All patients were age \(\geq 18\) years old and provided written informed consent at inclusion. Patients were assessed at diagnosis of breast cancer, and follow-up data were available through year 4 (Y4) after diagnosis at the time of the present analysis. Patients could choose to enrol in concomitant observational or interventional studies at any time during the follow-up period. CANTO study procedures were previously published.\(^17\) We used the Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) checklist for observational studies\(^18\) to assess the quality of our report.

**Study cohort**

For this analysis, we had access to information from 9597 patients with early breast cancer from the CANTO cohort. We excluded 81 patients with unknown participation status in concomitant research studies and those who were enrolled in concomitant studies but study information was unavailable or inaccessible. Overall, 9516 patients were included in the final analytic cohort (Figure 1). The aim of this study was to find associations between baseline cohort characteristics (collected at diagnosis of breast cancer) and likelihood of subsequent enrolment in clinical trials. Particularly, we assessed whether patients were enrolled in a clinical trial at any time over the course of their participation to the CANTO cohort study.

**Variables of interest**

**Clinical trials enrolment variables.** Clinical trials enrolment variables were: (i) clinical trials enrolment rate [e.g. whether patients enrolled or not in a clinical trial over the course of their participation in the CANTO study (from breast cancer diagnosis through Y4 after diagnosis)]; (ii) the number, and (iii) the type of clinical trial in which patients enrolled. Enrolment in a clinical trial was defined as participation in a concomitant interventional study comparing a new therapeutic approach to the standard of care, or testing a new clinical strategy through a direct intervention on the patient.\(^19\) Being included only in observational studies did not account as enrolment in clinical trials.

**Other variables**

**Clinical variables.** Clinical, sociodemographic and socioeconomic (educational level, income, and occupation), tumour and treatment-related characteristics, and data regarding cancer centre of care [including geographical region and patient volume (expressed as the number of patients included in the CANTO cohort by centre)] were available at breast cancer diagnosis and categorized as in Table 1. Anxiety and depression were assessed through the Hospital Anxiety and Depression Scale.\(^20\)

**Patient-reported outcomes.** Patient-reported outcomes (PROs) were collected using the European Organisation for
Research and Treatment of Cancer (EORTC) quality-of-life (QoL) questionnaires QLQ-C30,²¹ BR23,²² and FA12.²³ A higher-order QoL metric, the C30 Summary Score (C30 SumScore), was used to assess overall QoL. The C30 SumScore incorporates 13 (functional: physical, emotional, social, cognitive, and role; symptoms: fatigue, pain, nausea/vomiting, sleep disturbance, dyspnea, appetite loss, constipation, diarrhoea) of the 15 domains from EORTC QLQ-C30 (global health status and financial difficulties excluded).²⁴,²⁵ PROs were available in CANTO at diagnosis, and at Y1, Y2, and Y4 after diagnosis.

Survival outcomes. (i) A distant disease event (dDE) was defined as a distant breast cancer recurrence or death by any cause; (ii) invasive disease event (iDE) was defined as any invasive breast cancer recurrence (ipsilateral, regional, contralateral, or distant), second primary invasive cancer, or death by any cause; (iii) death by any cause (following standard DATECAN definitions).²⁶

Statistical analysis
Primary analyses. First, we used descriptive statistics to summarize clinical trials enrolment variables. Then, cohort characteristics at diagnosis were tabulated, overall and by enrolment in clinical trials. Chi-square tests and Wilcoxon rank sum tests were used to compare characteristics according to enrolment for categorical and continuous variables, as appropriate. A multivariable logistic regression model was used to assess factors associated with clinical trials enrolment at any time over the course of the participation to the CANTO cohort study. All patients who had information about enrolment status at any point over post-diagnosis follow-up were included.

Exploratory analyses. We assessed the relationship of enrolment in clinical trials with PROs and clinical outcomes. A propensity score matched analysis was carried out to reduce the potential influence of confounding factors and balance characteristics between patients enrolled and not enrolled in clinical trials. This propensity score analysis was carried out following standard guidelines previously reported in literature.²⁷ In order to calculate the propensity scores, we included in a logistic regression model all the variables assessed at baseline, including year of breast cancer diagnosis (as in Table 1), and we obtained the predicted probability of enrolment in clinical trials based on each patient characteristic. Once the propensity scores were calculated, subjects were matched by single score. Among patients selected and matched by propensity scores, the longitudinal evolution of PRO (EORTC QLQ-C30, BR23, and FA12) from diagnosis to Y4 was described according to clinical trials enrolment. A multiple linear regression model evaluated the association between clinical trials enrolment and C30 SumScore at Y4 after diagnosis, adjusting by C30 SumScore and covariates at baseline. The rates of dDE, iDE, and death by any cause were compared by enrolment in clinical trials using chi-square tests.

Sensitivity analyses. To confirm the robustness of our findings, we carried out the analyses of factors associated with enrolment among a subset of patients diagnosed with breast cancer from 2012 to 2015, who had complete follow-up available 4 years after diagnosis at the time of the most recent database update (n = 7810).

All tests were two-sided with a 95% confidence interval (CI) and a P value of <0.05 was considered statistically significant. Statistical analysis was carried out using SAS statistical software Version 9.4 (University of Bordeaux, Bordeaux, France).

RESULTS
Cohort characteristics
Patients’ characteristics are reported in Table 1. Overall, median age at diagnosis was 57.0 years [interquartile range (Q1-Q3) 17.0 years]. The majority of patients included in the cohort were postmenopausal women (62.8%), with a Charlson comorbidity score of 0 (80.8%), and reported at least doubtful symptoms of anxiety (60.5%). More than half of patients were
| Characteristic                          | Whole cohort | According to clinical trial enrolment |
|----------------------------------------|--------------|--------------------------------------|
|                                        | N (%         | Yes (%) 1716 (18.0) | No (%) 7800 (82.0) |
| **Baseline cohort characteristics**    |              |                       |                      |
| General characteristics                |              |                       |                      |
| Age at diagnosis, years                |              |                       |                      |
| Median (Q1-Q3)                         | 57.0 (48.5-65.5) | 57.7 (48.1-66.3) | 56.8 (48.6-65.4) | 0.1080 |
| Missing                                |              |                       |                      |
| BMI at diagnosis                       |              |                       |                      |
| Median (Q1-Q3)                         | 24.8 (22.0-28.7) | 25.3 (22.3-29.1) | 24.7 (21.9-28.7) | 0.0012 |
| Missing                                | 55           | 4                     | 51                   |
| Menopausal status                      |              |                       |                      |
| Premenopausal                          |              |                       |                      |
| Postmenopausal                         |              |                       |                      |
| Missing                                | 77           | 10                    | 67                   |
| Charlson comorbidity index             |              |                       |                      |
| 0                                     | 7049 (80.8)  | 1202 (17.0)           | 5849 (83.0)          | 0.0210 |
| 1+                                    | 1672 (19.2)  | 325 (19.4)            | 1347 (80.6)          | 0.3774 |
| Missing                                | 795          | 189                   | 606                  |
| Depression, categorical                |              |                       |                      |
| Normal                                 | 7087 (81.6)  | 1301 (18.4)           | 5786 (81.6)          | 0.3774 |
| Doubtful case/case                    |              |                       |                      |
| Missing                                | 827          | 136                   | 691                  |
| HADS depression score*                 |              |                       |                      |
| Mean (SD)                              | 4.2 (3.7)    | 4.1 (3.6)             | 4.3 (3.7)            | 0.0647 |
| Missing                                | 827          | 136                   | 691                  |
| Anxiety, categorical                   |              |                       |                      |
| Normal                                 | 3434 (39.5)  | 661 (19.2)            | 2773 (80.8)          | 0.0394 |
| Doubtful case/case                    |              |                       |                      |
| Missing                                | 832          | 136                   | 696                  |
| HADS anxiety score*                    |              |                       |                      |
| Mean (SD)                              | 9.0 (4.2)    | 8.7 (4.2)             | 9.0 (4.2)            | 0.0205 |
| Missing                                | 832          | 136                   | 696                  |
| Marital status                         |              |                       |                      |
| In a relationship                      | 6786 (77.9)  | 1236 (18.2)           | 5550 (81.8)          | 0.8015 |
| Not in a relationship                  | 1928 (22.1)  | 356 (18.5)            | 1572 (81.5)          | 0.8191 |
| Missing                                | 802          | 124                   | 678                  |
| Having a child                         |              |                       |                      |
| Yes                                    | 8313 (96.5)  | 1524 (18.3)           | 6789 (81.7)          | 0.8324 |
| No                                     | 303 (3.5)    | 57 (18.8)             | 246 (81.2)           | 0.5624 |
| Missing                                | 900          | 135                   | 765                  |
| Familiar history of breast cancer      |              |                       |                      |
| Positive                               | 2122 (22.7)  | 379 (17.9)            | 1743 (82.1)          | 0.8419 |
| Negative                               | 7230 (77.3)  | 1305 (18.0)           | 5925 (82.0)          | 0.8191 |
| Missing                                | 164          | 32                    | 132                  |
| Personal history of previous cancer    |              |                       |                      |
| Yes                                    | 735 (7.9)    | 118 (16.0)            | 617 (84.0)           | 0.1607 |
| No                                     | 8613 (92.1)  | 1561 (18.1)           | 7052 (81.9)          | 0.1607 |
| Missing                                | 168          | 37                    | 131                  |
| Previous breast surgery                |              |                       |                      |
| Yes                                    | 694 (7.3)    | 94 (13.5)             | 600 (86.5)           | 0.0014 |
| No                                     | 8776 (92.7)  | 1615 (18.4)           | 7161 (81.6)          | 0.8191 |
| Missing                                | 46           | 7                     | 39                   |
| Highest education level                |              |                       |                      |
| Primary or lower                       | 1285 (14.8)  | 255 (19.8)            | 1030 (80.2)          | 0.0605 |
| College graduate or higher             |              |                       |                      |
| Missing                                | 861          | 139                   | 722                  |
| Monthly household income (euros/month) |              |                       |                      |
| <2000                                  | 2288 (28.6)  | 430 (18.8)            | 1858 (81.2)          | 0.0763 |
| 2000-4000                              | 3730 (46.6)  | 710 (19.0)            | 3020 (81.0)          | 0.0014 |
| >4000                                  | 1590 (24.8)  | 332 (16.7)            | 1658 (83.3)          | 0.0014 |
| Missing                                | 1508         | 244                   | 1264                 |

Table 1. Baseline cohort characteristics, overall and according to clinical trials enrolment
diagnosed with stage II (41.6%) or III (9.7%) breast cancer, 76.0% of them had a hormone receptor-positive, human epidermal growth factor receptor 2-negative breast cancer, 81.0% received hormonal therapy, and 91.2% radiotherapy.

Enrolment in clinical trials and associated factors

The rate of enrolment in clinical trials was 18.0% in the overall cohort (1716/9516 patients); 89% of patients were recruited in only one clinical trial, 10% concomitantly in two

| N (%) | Whole cohort | According to clinical trial enrolment | Yes | No | P*
|---|---|---|---|---|---|
| Occupational class | | | | | |
| Chief executives, managers, and intellectual professionals | 3925 (45.1) | 668 (17.0) | 3257 (83.0) | 0.0020 |
| Middle-class workers\(^a\) | 3249 (37.4) | 612 (18.8) | 2637 (81.2) | |
| Self employed and manual workers\(^b\) | 3165 (35.7) | 270 (8.6) | 2895 (91.4) | |
| Unemployed, retired | 160 (1.8) | 43 (26.9) | 117 (73.1) | |
| Missing | 817 | 123 | 694 | |
| Region of residency | | | | | |
| Île-de-France | 2632 (27.7) | 391 (14.9) | 2241 (85.1) | <0.0001 |
| Centre/North of France\(^d\) | 4989 (52.4) | 940 (18.8) | 4049 (81.2) | |
| South of France\(^e\) | 1895 (19.9) | 385 (20.3) | 1510 (79.7) | |
| Missing | — | — | — | |
| Anonymized COC | | | | | |
| High volume | 2320 (24.4) | 373 (16.1) | 1947 (83.9) | 0.0004 |
| Intermediate volume | 6454 (67.8) | 1232 (19.1) | 5222 (80.9) | |
| Low volume | 742 (7.8) | 111 (15.0) | 631 (85.0) | |
| Missing | — | — | — | |
| COC and patients region of residence | | | | | |
| COC in the same region | 8591 (90.3) | 1550 (18.0) | 7041 (82.1) | 0.9424 |
| COC in a different region/foreign patients | 925 (9.7) | 166 (17.9) | 759 (82.1) | |
| Missing | — | — | — | |
| Tumour stage | | | | | |
| I | 4582 (48.7) | 644 (14.0) | 3938 (86.0) | <0.0001 |
| II | 3911 (41.6) | 864 (22.1) | 3047 (77.9) | |
| III | 912 (9.7) | 188 (20.6) | 724 (79.4) | |
| Missing | 111 | 20 | 91 | |
| Tumour subtype | | | | | |
| HR\(^+\)/HER2\(^-\) | 7186 (76.0) | 1318 (18.3) | 5868 (81.7) | 0.0951 |
| HR\(^-\)/HER2\(^+\) | 1317 (13.9) | 248 (18.8) | 1069 (81.2) | |
| HR\(^-\)/HER2\(^-\) | 954 (10.1) | 149 (15.6) | 805 (84.4) | |
| Missing | 59 | 1 | 58 | |
| Chemotherapy | | | | | |
| Yes | 5041 (53.0) | 997 (19.8) | 4044 (80.2) | <0.0001 |
| No | 4475 (47.0) | 719 (16.1) | 3766 (83.9) | |
| Missing | — | — | — | |
| Surgery | | | | | |
| Mastectomy | 2539 (26.7) | 456 (18.0) | 2083 (82.0) | 0.9100 |
| Conservative surgery | 6971 (73.3) | 1259 (18.1) | 5712 (81.9) | |
| Missing | 6 | 1 | 5 | |
| Hormone therapy | | | | | |
| Yes | 7679 (81.0) | 1442 (18.8) | 6237 (81.2) | 0.0003 |
| No | 1797\(^f\) (19.0) | 272 (15.1) | 1525 (84.9) | |
| Missing | 40 | 2 | 38 | |
| Radiotherapy | | | | | |
| Yes | 8661 (91.2) | 1630 (18.8) | 7031 (81.2) | <0.0001 |
| No | 831 (8.8) | 82 (9.9) | 749 (90.1) | |
| Missing | 24 | 4 | 20 | |
| Anti-HER2 therapy | | | | | |
| Yes | 1158 (12.2) | 230 (19.9) | 928 (80.1) | 0.0879 |
| No | 8341 (87.8) | 1485 (17.8) | 6856 (82.2) | |
| Missing | 17 | 1 | 16 | |

BMI, body mass index; COC, centre of care; HADS, Hospital Anxiety and Depression Scale; HER2, human epidermal growth factor receptor 2; HR, hormone receptor; Q1-Q3, interquartile range; SD, standard deviation.

\(^a\)Anxiety and depression were assessed using the Hospital Anxiety and Depression Scale (HADS).\(^20\)
\(^b\)Includes clerks, service and sales workers, technicians, and associates.
\(^c\)Includes farmers, workers, freelancers, artisans, and merchants.
\(^d\)Centre/North of France includes Brittany, Normandy, Hauts de France, Pays de la Loire, Grand Est, Bourgogne-Franche-Comté, Centre-Val de Loire, and other regions or foreign countries.
\(^e\)South of France includes Provence-Alpes-Côte d’Azur, Occitane, Corsica, Nouvelle Aquitaine, and Auvergne-Rhône-Alpes.
\(^f\)Some 87.3% had an HR\(^+\)/HER2\(^-\) early breast cancer and 12.7% an HR\(^+\)/HER2\(^+\) early breast cancer.
\(^g\)Some 6.6% had an HR\(^+\)/HER2\(^-\) early breast cancer, 29.2% an HR\(^+\)/HER2\(^+\) early breast cancer, and 95.2% a triple-negative breast cancer.

*Chi square for categorical variables, Wilcoxon rank sum test for continuous variables.
clinical trials, whereas ~1% were recruited in three or four clinical trials. The majority of patients were enrolled in phase III drug evaluation studies (n = 641), phase III not-drug evaluation studies (n = 549), and phase II drug evaluation studies (n = 398), whereas only a few patients (n = 59) were recruited in early-phase clinical trials (Table 2).

After multivariable adjustment, factors independently associated with enrolment in clinical trials were: age at diagnosis [40-50 versus >65 years, odds ratio (OR) 0.79 (95% CI 0.64-0.98), P = 0.0300; 50-65 versus >65 years, OR 0.75 (95% CI 0.63-0.90), P = 0.0021], not reporting anxiety symptoms at diagnosis [normal versus doubtful case/case OR 1.17 (95% CI 1.01-1.35), P = 0.0363], region of residency [centre/North versus Ile de France OR 1.26 (95% CI 1.05-1.51) P = 0.0145; South versus Ile de France OR 1.51 (95% CI 1.20-1.89) P = 0.0004], being followed up in an intermediate volume centre of care [intermediate versus low volume centre OR 1.43 (95% CI 1.07-1.92), P = 0.0161] tumour stage [stage II versus stage I OR 1.56 (95% CI 1.33-1.84), P < 0.0001; stage III versus stage I OR 1.32 [95% CI 1.01-1.73], P = 0.0451], and adjuvant treatments such as hormonal therapy [yes versus no OR 1.34 (95% CI 1.11-1.62), P = 0.0021] and radiotherapy [yes versus no OR 1.98 (95% CI 1.43-2.76), P < 0.0001]. Full results are displayed in Table 3 and Supplementary Figure S2, available at https://doi.org/10.1016/j.esmoop.2022.100513; details regarding rates of enrolment in clinical trials by different regions are displayed in Supplementary Figure S3, available at https://doi.org/10.1016/j.esmoop.2022.100513.

Exploratory analyses: PROs and survival outcomes by enrolment in clinical trials

The propensity scores matching analysis led to the selection of 2118 patients with similar propensity scores. Baseline cohort characteristics of the propensity score selected population are displayed in Supplementary Table S1, available at https://doi.org/10.1016/j.esmoop.2022.100513. C30 SumScore behaved similarly in patients enrolled and not enrolled in a clinical trial (Figure 2). Scores for all QLQ-C30, BR23, and FA12 domains at diagnosis, Y1, Y2, and Y4 are displayed in Supplementary Tables S2 and S3, available at https://doi.org/10.1016/j.esmoop.2022.100513. There was a significant association between the C30 SumScore at Y4 and the enrolment in clinical trials (mean difference versus not enrolled 1.37, 95% CI 0.03-2.71, P = 0.0458) (Supplementary Table S4, available at https://doi.org/10.1016/j.esmoop.2022.100513). Among the propensity score selected patients, 7.3% of those enrolled in clinical trials had a dDE compared with 10.1% among those not enrolled (P = 0.0206). In addition, the rate of IDE was 8.2% among patients enrolled in clinical trials and 10.5% in those not enrolled (P = 0.0732). Lastly, death by any cause was 2.8% among patients enrolled in clinical trials and 3.7% among those not enrolled (P = 0.2707). The overall number of events by enrolment in clinical trials in the cohort of 2118 patients selected after the propensity score matched analysis is displayed in Supplementary Table S5, available at https://doi.org/10.1016/j.esmoop.2022.100513.

Sensitivity analyses

We carried out a sensitivity analysis on a subset of patients diagnosed with breast cancer from 2012 to 2015, who had complete follow-up available 4 years after diagnosis (n = 7810). Factors associated with enrolment in clinical trials were consistent with the main analysis. Results are displayed in Supplementary Table S6, available at https://doi.org/10.1016/j.esmoop.2022.100513.

DISCUSSION

Our analysis showed that among patients with early breast cancer, one in five enrolled in a clinical trial over a period of time extending from diagnosis to 4 years after, with the majority of them participating in phase III clinical trials and few patients participating in more than one study. We found some differences in clinical (age, stage of disease, anxiety) and geographical (region of residency, facilities of care) characteristics between patients enrolled in clinical trials and not. Trial participation did not seem to pose additional burden on patient-reported QoL 4 years after diagnosis, and seemed associated with improved clinical outcomes, as shown by similar PROs and lower rates of iDEs among patients who were enrolled in a clinical trial.

We believe that this study offers valuable insights into factors associated with clinical trial enrolment and expands upon the existing literature on the topic. First, we found that elderly women had the highest likelihood of enrolment in clinical trials in this cohort. Contrary to our findings, older adults are generally underrepresented in clinical trials. This was usually attributed to medical comorbidities, stringent trial eligibility criteria, and patient or physician misconceptions about the risks of enrolment among elderly individuals. A possible explanation of a relatively good representation of the >65 years of age category in this study is that many patients in our cohort could access trials focused on supportive care, symptom management, and amelioration of QoL such as, for example, the ART-THERAPIE trial or the BEAUTY trial. Factors contributing to the lower enrolment rates among younger patients, however, including those aged 40-65 years in our cohort, are less well

| Type of study               | No. patients enrolled | N = 1716 (%) |
|-----------------------------|-----------------------|--------------|
| Phase III drug assessing    | 641 (37.3)            |              |
| Phase III not drug assessing| 549 (32.0)            |              |
| Phase III supportive care   | 136 (7.9)             |              |
| Phase II-III                | 3 (0.2)               |              |
| Phase II-III supportive care| 19 (1.1)              |              |
| Phase II drug assessing     | 396 (23.2)            |              |
| Phase II not drug assessing | 20 (1.2)              |              |
| Phase II supportive care    | 21 (1.2)              |              |
| Phase I-I                   | 57 (3.3)              |              |
| Phase I                     | 2 (0.1)               |              |

*Numbers do not add up to 1716 as patients could be enrolled in more than 1 type of clinical trial at the same time. Note, among patients enrolled in clinical trials, there were 90 patients who also participated in concomitant observational studies.
understood, including in previous literature. Among these, the physician’s failure to propose access to innovation was described as a strong barrier to enrolment in younger patients in previous studies.15,32,33

Second, some patient-related clinical factors were associated with enrolment. Particularly, an anxiety trait was associated with lower enrolment in clinical trials. As also suggested by previous studies, uncertainties, fear of potential adverse events, fear of delay of initiation of antineoplastic drugs, fear of being enrolled in the placebo arm due to randomization procedures, or possible painful procedures, may possibly reduce willingness to participate in clinical studies.34 For instance, a survey by Mancini et al.35 on 115 women enrolled in clinical trials from 21 centres in France pointed out how 43.0% of participants expressed mild regret to have chosen to participate, and 25.8% expressed moderate to strong regret after agreeing to participate in a randomized, controlled clinical trial.35 On the contrary, a strong clinical barrier to the participation in clinical trials previously reported in the literature, such as comorbidities,36 was not associated with enrolment in our cohort.

Third, we did not find major differences in enrolment rates across socioeconomic strata in our cohort, as income and education level did not seem to be associated with enrolment. We acknowledge, however, that these social determinants may represent important drivers of enrolment in health care systems other than the French system.

Financial issues were identified as an important barrier to clinical trial participation, especially in the United States,37 where insurances may not cover non-routine services.38 Whereas in France many health services are often accessible with no extra cost to the whole population, due to universal health coverage, in some countries, such as the United States, patients may have to deal with additional costs when participating in clinical trials, especially when this requires them to travel to their follow-up centres for additional tests or procedures that are only available locally, or cannot be obtained elsewhere.39

Fourth, centre-related features seemed to be one of the most relevant determinants of enrolment in clinical trials. This finding may relate to availability of research infrastructures that can facilitate enrolment, both for patients and for physicians. Facilitators of enrolment may indeed come from the improvement of local research infrastructures, providing facilities to physicians that can optimize the process of recruitment and trial managing.

In addition, we carried out exploratory analyses that also yielded interesting findings. We showed that enrolment in clinical trials seems to be associated with better PRO 4 years after diagnosis and lower recurrence rates. Whether the enrolment in clinical trials improves survival beside the clinical setting or trial outcomes still remains debated. Previously,
one study by Chow et al. proved that enrolment into cancer trials could independently predict lower overall and cancer-specific mortality in patients with common cancers. Although exploratory, results of our study are consistent with these findings and provide further insight into the relationship between enrolment in trials and relapse among a specific population, such as patients with early breast cancer.

We recognize that our study has some limitations. Our cohort is selected from patients that participated in CANTO, which is per se a multicenter cohort study. Indeed, the high proportion of comprehensive cancer centres in the recruiting institutions might have led to an overestimation of the results. CANTO focuses on women diagnosed with breast cancer and followed up in France, therefore this analysis may not be generalizable to all women and health care systems of other countries. Some attrition was present with increasing missing data, particularly in PROs at later time points. CANTO did not include information about the date and duration of enrolment clinical trials, as participation could happen anytime over 4 years after diagnosis. To accommodate for the potential of premature study termination due to breast cancer recurrence or death, however, all models included breast cancer prognostic factors. CANTO includes only patients with stage I-II-III breast cancer without evidence of active disease or relapse, therefore we acknowledge potential bias from study termination for patients who develop disease progression. Analyses of clinical outcomes may be underpowered due to the limited number of events in this early-stage cohort of women with breast cancer and limited follow-up time for survival events. For these reasons, survival outcomes were summarized descriptively, as opposed to being assessed in a formal time-to-event analysis by enrolment group, and our results should be interpreted with caution as they were considered exploratory. Strengths include that this is the first multicentre prospective study to focus exclusively on the enrolment in clinical trials among women diagnosed with early breast cancer, with availability of a large amount of longitudinal PRO up to 4 years after diagnosis.

**Conclusions**

This study focused on the enrolment rate and factors related to enrolment in clinical trials among women diagnosed with early breast cancer. One in five women were enrolled in a clinical trial after diagnosis of early-stage breast cancer. In this population, the rate of enrolment exceeded what was previously found in different settings. Whereas patients seem adequately represented irrespective of clinical and treatment-related features, mostly clinical (age, anxiety traits) and geographical (region of residency, facilities of care) factors seem to be those associated with access to innovation in clinical trials. Enrolment did not negatively impact QoL and there were suggestions of improved clinical outcomes among women who participated in clinical trials. Access to innovation should be facilitated and encouraged by overcoming factors associated with impaired recruitment.

**FUNDING**

This work was supported by the French Government under the ‘Investment for the Future’ program managed by the National Research Agency (ANR) [grant number ANR-10-COHO-0004]. Antonio Di Meglio is supported by a Career Pathway Grant in Symptom Management from Conquer Cancer, the ASCO
Foundation and Rising Tide Foundation for Clinical Cancer Research. This study was supported by grants from Odyssea (no grant number), and Foundation Gustave Roussy (no grant number).

DISCLOSURE
BP reports personal fees from AstraZeneca, Myriad, Pfizer, Pierre Fabre, Daichi Sankyo, and non-financial support from Daichi Sankyo, Merus, Puma, Pfizer, AstraZeneca. All other authors have declared no conflicts of interest.

REFERENCES

1. Braunholtz DA, Edwards SJL, Lilford RJ. Are randomized clinical trials good for us (in the short term)? Evidence for a “trial effect.”. J Clin Epidemiol. 2001;54(3):217-224.

2. Shahar T, Nossek E, Steinberg DM, et al. The impact of enrollment in clinical trials on survival of patients with glioblastoma. J Clin Neurosci. 2012;19(11):1530-1534.

3. Chow CJ, Habermann EB, Abraham A, et al. Does enrollment in cancer trials improve survival? J Am Coll Surg. 2013;216(4):774-781.

4. Arrieta O, Carmona A, Ramirez-Tirado LA, et al. Survival of patients with advanced non-small cell lung cancer enrolled in clinical trials. Oncology. 2016;91(4):185-193.

5. National Comprehensive Cancer Network. NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines). Available at NCCN.org. Accessed September 13, 2019.

6. Murthy VH, Krumholz HM, Gross CP. Participation in cancer clinical trials: race-, sex-, and age-based disparities. JAMA. 2004;291(22):2720-2726.

7. Carey M, Boyes AW, Smits R, Bryant J, Waller A, Oliver I. Access to clinical trials among oncology patients: results of a cross sectional survey. BMC Cancer. 2017;17(1):653.

8. U.S. Food and Drug Administration. 2015-2016 Global Participation In Trials Report. July 2017.

9. Castro EM, Van Regenmortel T, Vanhaeckt K, Sermeus W, Van Hecke A. Patient empowerment, patient participation and patient-centeredness in hospital care: a concept analysis based on a literature review. Patient Educ Couns. 2016;99(12):1923-1939.

10. Unger JM, Vaidya R, Hershman DL, Minasian LM, Fleury ME. Systematic review and meta-analysis of the magnitude of structural, clinical, and physician and patient barriers to cancer clinical trial participation. J Natl Cancer Inst. 2019;111(3):245-255.

11. Unger JM, Cook E, Tai E, Bleyer A. The role of clinical trial participation in cancer research: barriers, evidence, and strategies. Am Soc Clin Oncol Educ Book. 2016;35:185-198.

12. Kincaid E. Advanced cancer treatments far from big-name hospitals. The Wall Street Journal. Available at https://www.healthleadersmedia.com/strategy/advanced-cancer-treatments-far-big-name-hospitals. Accessed March 6, 2017.

13. Mills EJ, Seely D, Rachlis B, et al. Barriers to participation in clinical trials of cancer: a meta-analysis and systematic review of patient-reported factors. Lancet Oncol. 2006;7(2):141-148.

14. Comis RL, Miller JD, Aldigé CR, Krebs L, Stoval E. Public attitudes toward participation in cancer clinical trials. J Clin Oncol. 2003;21(5):830-835.

15. Mazouni C, Deneuve J, Arnedos M, et al. Decision-making from multidisciplinary team meetings to the bedside: factors influencing the recruitment of breast cancer patients into clinical trials. Breast. 2014;23(2):170-174.

16. Besle S, Schultz E, Hollebecque A, et al. Organisational factors influencing early clinical trials enrollment: Gustave Roussy experience. Eur J Cancer. 2018;98:17-22.

17. Vaz-Luis I, Cottu P, Mesleard C, et al. UNICANCER: French prospective cohort study of treatment-related chronic toxicity in women with localised breast cancer (CANTO). ESMO Open. 2019;4(5):e000562-e000562.

18. von Elm E, Altman DG, Egger M, Pocock SJ, Gøtzsche PC, Vandenbroucke JP. The Strengthening the Reporting of Observational Studies in Epidemiology (STROBE) statement: guidelines for reporting observational studies. J Clin Epidemiol. 2008;61(4):344-349.

19. WHO International Clinical Trials Registry Platform. 2021. Available at https://www.who.int/clinical-trials-registry-platform. Accessed April 20, 2022.

20. Zigmund AS, Snaith RP. The hospital anxiety and depression scale. Acta Psychiatr Scand. 1983;67(6):361-370.

21. Aaronson NK, Ahmedzai S, Bergman B, et al. The European Organization for Research and Treatment of Cancer QLQ-C30: a quality-of-life instrument for use in international clinical trials in oncology. J Natl Cancer Inst. 1993;85(5):365-376.

22. Sprangers MA, Groenvold M, Arraras JI, et al. The European Organization for Research and Treatment of Cancer breast cancer-specific quality-of-life questionnaire module: first results from a three-country field study. J Clin Oncol. 1996;14(10):2756-2768.

23. Weis J, Tomaszewski KA, Hammerlid E, et al. International psychometric validation of an EORTC quality of life module measuring cancer related fatigue (EORTC QLQ-FA12). J Natl Cancer Inst. 2017;109(5).

24. Giesinger JM, Kieffer JM, Fayers PM, et al. Replication and validation of higher order models demonstrated that a summary score for the EORTC QLQ-C30 is robust. J Clin Epidemiol. 2016;69:79-88.

25. Husson O, de Rooij BH, Kieffer J, et al. The EORTC QLQ-C30 summary score as prognostic factor for survival of patients with cancer in the “Real-World”: results from the population-based PROFILES registry. Oncologist. 2020;25(4):e722-e732.

26. Gourgou-Bourdaise S, Cameron D, Poortmans P, et al. Guidelines for time-to-event end point definitions in breast cancer trials: results of the DATECAN initiative (Definition for the Assessment of Time-to-event Endpoints in CANcer trials). Ann Oncol. 2015;26(5):873-879.

27. Yao Xi, Wang X, Speicher PJ, et al. Reporting and guidelines in propensity score analysis: a systematic review of cancer and cancer surgical studies. J Natl Cancer Inst. 2017;109(8):dixw233.

28. Kemeny MM, Peterson BL, Kornblith AB, et al. Barriers to clinical trial participation by older women with breast cancer. J Clin Oncol. 2009;27(11):2269-2275.

29. Denson AC, Mahjali A. Participation of the elderly population in clinical trials: barriers and solutions. Cancer Control. 2014;21(3):209-214.

30. Joly F, Pasquier D, Hanzen C, et al. Impact of art therapy (AT) on fatigue and quality of life (QoL) during adjuvant external beam irradiation (EBI) in breast cancer patients (pts): a randomized trial. Ann Oncol. 2016;27(suppl 6):vi499.

31. Saghatcian M, Lucas B, Charles C, et al. BEAUTY and the breast: is adjuvant chemotherapy the right time for a beauty boost? Lessons learned from a large randomized controlled trial. Qual life Res. 2022;31:723-732.

32. Kim ES, Bruinooge SS, Roberts S, et al. Broadening eligibility criteria to make clinical trials more representative: American Society of Clinical Oncology and Friends of Cancer Research Joint Research statement. J Clin Oncol. 2017;35(33):3737-3744.

33. Siembida EJ, Loomans-Kropp HA, Trivedi N, et al. Systematic review of barriers and facilitators to clinical trial enrollment among adolescents and young adults with cancer: identifying opportunities for intervention. Cancer. 2020;126(5):949-957.

34. Almeida L, Kashdan TB, Nunes T, Coelho R, Albino-Teixeira A, Soares-da-Silva P. Who volunteers for phase I clinical trials? In Decision making. J Clin Epidemiol. 2012;65(6):635-642.

35. Zaorsky NG, Zhang Y, Walter V, Tchelebi LT, Chinchilli VM, Gusani NJ. Prevention and management of chemotherapy-related toxicities in women with breast cancer (CANTO). J Oncol Pract. 2013;9(5):251-257.

36. Nipp RD, Powell E, Chabner B, Moy B. Recognizing the financial burden of cancer patients in clinical trials. Oncologist. 2015;20(6):572-575.