Randomised, open-label, multicentric phase III trial to evaluate the safety and efficacy of palbociclib in combination with endocrine therapy, guided by ESR1 mutation monitoring in oestrogen receptor-positive, HER2-negative metastatic breast cancer patients: study design of PADA-1

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ABSTRACT
Introduction The combination of a CDK4/6 inhibitor with an aromatase inhibitor (AI) has recently become the gold standard for AI-sensitive first line treatment of oestrogen receptor-positive (ER+) HER2-negative (HER2−) advanced breast cancer. However, most patients receiving this combination will ultimately progress and require further therapies. Several studies have demonstrated that the onset of an ESR1 gene mutation lead to AlS resistance in the advanced setting. ESR1 mutations can be detected in circulating tumour DNA (ctDNA) using a digital PCR assay. Our study aims to prove the clinical efficacy of periodic monitoring for emerging or rise of ESR1 mutations in ctDNA to trigger an early change from AI plus palbociclib to fulvestrant plus palbociclib treatment while assessing global safety.

Methods PADA-1 is a randomised, open-label, multicentric, phase III trial conducted in patients receiving AI and palbociclib as first line therapy for metastatic ER+HER2- breast cancer. 1000 patients will be included and treated with palbociclib in combination with an AI. Patients will be screened for circulating blood ESR1 mutation detection at regular intervals. Patients for whom a rising circulating ESR1 mutation is detected without tumour progression (up to N=200) will be randomised (1:1) between (1) Arm A: no modification of therapy; and (2) Arm B: palbociclib in combination with fulvestrant, a selective ER down-regulator. At tumour progression, an optional crossover will be offered to patients randomised in arm A. The coprimary endpoints are (1) Grade ≥3 haematological toxicities and their associations with baseline characteristics and (2) progression-free survival in randomised patients.

Strengths and limitations of this study
► PADA-1 is the first-of-its-kind trial to engage the ‘subclone war’, by targeting ESR1 mutation right after they appear and before disease progression.
► Because of its large size and design, if positive, PADA-1 could be practice changing.
► More potent oral selective oestrogen receptor degraders could yield to greater benefit than fulvestrant.
► Cross-over was optional and postprogression treatments are not standardised, limiting the power of secondary efficacy endpoints that involve postprogresion therapies.
► No hypothesis is made to conclude about the toxicity of the palbociclib+fulvestrant combination.

BACKGROUND
Significant progresses have been made in detecting and treating breast cancer over the last three decades. Although the relative mortality linked to breast cancer is declining,
around 20% of patients with initial early breast cancer will ultimately develop metastases despite adequate adjuvant treatments. Furthermore, 5%–7% of breast cancer patients present with metastases at diagnosis in Western countries. Even though survival expectancy is improving, in most cases, breast cancer is considered incurable once it reaches the metastatic stage.\(^1\) This study investigates the first-line therapy of the most frequent breast cancer subtype oestrogen receptor-positive (ER+) HER2-negative (HER2−) metastatic breast cancer (about 70% of metastatic breast cancer cases).

In patients diagnosed with metastatic or locally advanced ER+HER2− breast cancer not amenable to local curative treatments (surgery or radiotherapy), systemic therapy must be initiated to stop and eventually reduce the tumour burden and increase the patient life expectancy. Costs and treatment-related toxicity are limited in patients receiving endocrine therapy, prompting conferences and cancer guidelines (ABC breast cancer consensus conference; National Cancer Center Network clinical practice guidelines) to recognise endocrine therapy as the gold standard for first-line therapy of metastatic ER+HER2− breast cancer for more than 10 years.\(^2^3\)

Once tumour progression has been observed under first-line therapy, there are no guidelines for the choice of subsequent lines of treatments. However, some considerations apply: endocrine therapy should be preferred unless chemotherapy seems to be a better option—because of poor general status, ‘visceral crisis’ (that can be defined, according to ABC Consensus, as severe organ dysfunction assessed by signs and symptoms, laboratory studies, and rapid disease progression).

Former trials presented here by chronological order, used the same palbociclib regimen (125 mg/day orally for 3 weeks followed by 1 week off) and dose adaptation than in PADA-1 trial. In the randomised phase II study PALOMA-1, first-line treatment with the combination of palbociclib plus letrozole achieved a progression-free survival (PFS) of 20.2 months vs 10.2 months with letrozole alone (p=0.0004).\(^4\) Best overall response rate and clinical benefit rate (complete/partial response plus stable disease) for the combination vs letrozole alone were 43% vs 33% and 81% vs 58%, respectively. Subgroup analysis showed that the PFS benefit for the combination was consistent across all subgroups including age, sites of metastatic disease and previous adjuvant therapies. The most common adverse reaction in patients receiving palbociclib plus letrozole was neutropenia. Based on this phase II study, the FDA granted in 2015 accelerated approval to palbociclib for use in combination with aromatase inhibitor (AI) for the treatment of postmenopausal women with HR+HER2− metastatic breast cancer as initial endocrine-based therapy for their metastatic disease. Before granting approval as first-line therapy, the EMA chose to wait for the results of the first-line phase III trial, PALOMA-2 (see beyond).

The PALOMA-3 trial assessed the efficacy of palbociclib and fulvestrant in endocrine-resistant advanced breast cancer.\(^5\) In this study, 521 patients with HR+HER2−advanced metastatic breast cancer whose cancer had relapsed or progressed on prior endocrine therapy were randomised to palbociclib and fulvestrant or placebo and fulvestrant. Premenopausal and perimenopausal women were allowed in this study and received Luteinising hormone-releasing hormone agonist. The median PFS was 9.2 months (95% CI 7.5 to not estimable) with palbociclib-fulvestrant and 3.8 months (95% CI 3.5 to 5.5) with placebo-fulvestrant (HR 0.42; 95% CI 0.32 to 0.56; p<0.001). The most common grade 3 or 4 adverse events (AEs) in the palbociclib-fulvestrant group were neutropenia (62.0%, vs 0.6% in the placebo-fulvestrant group), leucopenia (25.2% vs 0.6%), anaemia (2.6% vs 1.7%), thrombocytopenia (2.3% vs 0%), and fatigue (2.0% vs 1.2%). Febrile neutropenia was reported in only 0.6% of palbociclib-treated patients and 0.6% of placebo-treated patients. Based on the PALOMA-3 study, palbociclib was approved in Europe and made available in France for the treatment of advanced, endocrine therapy-resistant HR+HER2− metastatic breast cancer.

In the PALOMA-2 trial, 666 postmenopausal patients with no prior systemic therapy for metastatic breast cancer were randomised 2:1 to receive palbociclib (same schedule than in the above-mentioned studies) combined with an AI (letrozole) or placebo +letrozole until disease progression, consent withdrawal or death. In that study, median PFS was 24.8 months (palbociclib +letrozole) vs 14.5 months (placebo +letrozole) (HR=0.58; 95% CI 0.46 to 0.72; p=0.000001). Subgroup analyses regarding the efficacy were negative. Overall response rate was improved with palbociclib +letrozole (42.1% vs 34.7%; p=0.03; 55.3% vs 44.4% in patients with measurable disease; p=0.01). Clinical benefit rate (including patients with stable disease) was 84.9% vs 70.3% (p<0.0001).\(^6\) The first biomarker analysis presented at the 2016 ESMO meeting could not identify predictive markers of palbociclib efficacy.\(^7\) Common AEs (all grades) observed in the PALOMA-2 study with palbociclib +letrozole vs placebo +letrozole were neutropenia (79.5% vs 63.5%), fatigue (37.4% vs 27.5%), nausea (35.1% vs 26.1%), arthralgia (33.3% vs 33.8%) and alopecia (32.9% vs 15.8%). Grade 4 neutropenia was observed in 10% of patients. Febrile neutropenia was only seen with palbociclib and letrozole (1.8% of patients). Permanent discontinuation due to AEs was 9.7% (palbociclib +letrozole) vs 5.9% (placebo +letrozole). Overall survival data were immature.

Palbociclib is the first-in-class CDK4/6 inhibitor and other inhibitors are being developed. In October 2016, the results of a pivotal phase III trial testing another CDK4/6 inhibitor have been disclosed at the ESMO congress. In the MONALEESA-2 trial,\(^8\) the addition of ribociclib to letrozole significantly improved PFS in postmenopausal women with ER+advanced breast cancer, with a HR of 0.56. Median PFS, response rates, and toxicities (except for a possibly higher rate of grade 3/4 liver toxicity in MONALEESA-2) reported in both study arms were alike.

those observed in the PALOMA-2 trial. Palbociclib can be prescribed in the USA as first-line therapy since the results of PALOMA-1 and became the standard of care in first line, with more than 40,000 US patients treated as of November 2016. In Europe, on 10 November 2016, the European Commission (EC) has approved palbociclib (IBRANCE) in combination with an AI for the treatment of women with HR +HER2- locally advanced or metastatic breast cancer. The approval also covers the use of IBRANCE palbociclib was also approved in combination with fulvestrant for women who have received prior endocrine therapy (as proposed in PADA-1).

ESR1 (oestrogen receptor 1) is the gene coding for the ERα protein. Several studies have shown that ESR1 mutations are rare (<1%) at the initial stage of cancer and occurred during the metastatic stage. Approximately 12 ESR1 point mutations have been described, with a hotspot in codons 536–538 in exon 8. These mutations result in ligand-independent ER activity. In vitro and preclinical data suggest that ESR1 mutations lead to AIs resistance. Schiavon et al. have developed multiplex droplet digital PCR (ddPCR) assays for ESR1 mutations in circulating tumour DNA (ctDNA) and investigated the clinical relevance and origin of ESR1 mutations in 171 advanced breast cancer women. ESR1 mutation status in ctDNA (see beyond) showed high concordance with contemporaneous tumour biopsies and was accurately assessed in samples shipped at room temperature in preservative tubes. ESR1 mutations were found exclusively in ER+ breast cancer patients previously exposed to AI. Patients with ESR1 mutations had a substantially shorter PFS on subsequent AI-based therapy (HR 3.1; 95% CI 1.9 to 23.1; p=0.004). In the SoFEA trial, ESR1 mutations were found in 39.1% of patients (63 of 161), of whom 49.1% (27 of 55) were polyclonal, with rates of mutation detection unaffected by delays in processing of archived plasma. Patients with ESR1 mutations had improved PFS with fulvestrant (n=45) compared with exemestane (n=18; HR 0.52; 95% CI 0.3 to 0.92; p=0.02), whereas wild-type ESR1 patients had similar PFS with either treatment (HR 1.07; 95% CI 0.68 to 1.67; p=0.77). In the PALOMA-3 trial (advanced metastatic breast cancer, see above), ESR1 mutations were found in the plasma of 25.3% of patients (91 of 360), of whom 28.6% (26 of 91) were polyclonal, with mutations associated with acquired resistance to prior AI.11 Fulvestrant plus palbociclib improved PFS compared with fulvestrant plus placebo in both ESR1 mutant (HR 0.43; 95% CI 0.25 to 0.74; p=0.002) and ESR1 wild-type patients (HR 0.49; 95% CI 0.35 to 0.70; p<0.001). Therefore, ESR1 mutations can be robustly identified with ctDNA analysis and predict resistance to subsequent AI therapy. Thus, detection of ESR1-activating mutations may be relevant for guiding clinicians between endocrine and non-endocrine therapy.

Tumour cells release fragments of DNA in blood because of their high turnover (ctDNA) and, in cancer patients, ctDNA represents a variable fraction of cell-free DNA (cfDNA). ctDNA differ from normal cfDNA by the presence of cancer-related mutations, as ctDNA fragments harbour the genetic alterations found in the tumour cells of origin. Indeed, several reports showed high concordance between ctDNA mutations, when detectable, and matched tumour mutations.12-14 ctDNA fraction in blood is extremely variable, and can represent from 0.01% to more than 50% of cfDNA.15

Currently, ’liquid biopsy’ approaches depict the mutational landscape of given cancer by qualitative ctDNA analysis; however, the quantitative analysis of ctDNA ‘load’ in the blood may also be used as a dynamic marker to monitor tumour burden and response to therapy. Several studies reported a correlation between ctDNA changes during therapy with tumour sensitivity to treatment. ctDNA changes may be predictive markers of both responses at the start of treatment and secondary resistance during treatment. In addition, two reports showed that rising ctDNA levels can be detected several months before the onset of overt metastatic lesions during follow-up of patients treated for primary breast cancer. The median lead time between rising ctDNA levels and diagnosis of metastatic progression being around 10 months.21 In the present proposal, we will perform ctDNA analyses using state of the art platforms, with good sensitivity (ddPCR), as part of a collaborative work of the UNICANCER ctDNA group.

In summary, we hypothesised that early change of treatment from AI plus palbociclib to fulvestrant plus palbociclib based on rising ctDNA levels can improve prognosis of patient. To test the clinical utility of this strategy, we proposed a randomised, open-label, multicentric, phase III trial conducted in patients receiving AI and palbociclib as first-line therapy for metastatic ER+ HER2-negatve breast cancer.

METHODS/DESIGN

This study, including the option of cross-over (not mandatory), has been approved by the French medicines agency (ANSM, Agence Nationale de Sécurité du Médicament et des produits de santé) and by an ethics committee (Comité de Protection des Personnes Ouest IV- Nantes) in January 2017. The overall study design is shown in figure 1.

Aim of the study and endpoints

The coprimary objectives are (1) to assess whether a change of the endocrine therapy associated with palbociclib (namely, an early switch from AI with palbociclib to fulvestrant with palbociclib following ctDNA detection) will benefit patients in which rising ESR1 mutations are detected during treatment with palbociclib and AI, and (2) to assess the global safety of the combination of palbociclib +endocrine therapy in the whole population of patients, throughout the study, including the potential sequential administration of fulvestrant plus palbociclib, with focus on haematological toxicities. Therefore, coprimary endpoints are (1) PFS measured from the time of
Palbociclib and ctDNA for ESR1 mutation detection

Figure 1 Study scheme. AI, aromatase inhibitor; ctDNA, circulating tumour DNA; PADA-1, Palbociclib and ctDNA for ESR1 mutation detection.

randomisation (following rising ESR1 mutation detection) to the time of tumour progression (as assessed by the investigator per RECIST V.1.1 criteria\(^2\)) or death (whichever comes first)—in randomised patients. Patients who have not experienced disease progression and have not died will be censored at the date of last tumour evaluation. Efficacy analyses will be performed using the local radiologist’s/investigator’s tumour assessments as primary data source. (2) Safety assessed by collection of grade ≥3 AEs, using the Common Terminology Criteria for Adverse Events, V.4.03—in all included patients. The coprimary analysis will focus on grade ≥3 haematological toxicities and their associations with baseline characteristics.

Secondary and translational objectives and endpoints are listed in table 1.

Recruitment
During the first step, 1000 patients are planned to be included in 83 French centres. Based on the capacities of inclusion from each participating centre, the recruitment phase should last approximately 36 months.

Eligibility criteria are listed in box 1. After inclusion and non-inclusion criteria have been fulfilled and the patient consent has been obtained, the patient will be included in the step 1 treatment of the trial. Only patients with rising ESR1 mutations will be randomised in the step 2 treatment to either arm A to continue the same therapy (ie, palbociclib—AI combination) or arm B with an early switch to receive palbociclib in association with fulvestrant. Patients will be randomly assigned to one of the two arms (A or B) in a 1:1 ratio, using minimisation procedure\(^3\) stratified according to the presence or not of visceral involvement (defined as any lung, liver, pleura and brain metastasis) and time to rising ESR1 mutation detection (≥12 months vs <12 months; defined by the time between the date of the first ESR1 mutation detection sampling (baseline evaluation or cycle 1 day 1 visit) and the date of ESR1 mutation detection). The randomisation will be performed using the module of the eCRF/Ennov Clinical software V.7.5.40.7. After the consent form has been signed and all inclusion/non-inclusion criteria checked for step 2, the investigator will proceed with the randomisation through the R&D Unicancer online electronic case report form. Patients randomised in arm A will be offered to cross-over and be treated by fulvestrant + palbociclib, after disease progression under AI+palbociclib treatment, since Fribbens et al observed an improvement of PFS in patients treated this combination. Palbociclib will be continued as the same dose as before progression (ie, if palbociclib is reduced during step 2 for toxicity, it will not be increased again during step 3). For these patients, the third step (and the trial treatment) will end following tumour progression under fulvestrant + palbociclib.

Trial treatment
The investigational medicinal products are palbociclib in combination with the standard endocrine therapies, either AI (letrozole, anastrozole or exemestane) or an ER antagonist (fulvestrant). The endocrine therapies and palbociclib should be administered and dose adapted according to Summary of Products Characteristic recommendations and local practice. Palbociclib at a dose of 125 mg will be administered orally at the same time once a day for 21 days followed by 7 days off treatment of every 28-day cycle. Palbociclib capsules will be administered together with AI (letrozole, anastrozole or exemestane).
Table 1 Secondary and translational objectives and endpoints

<table>
<thead>
<tr>
<th>Secondary objectives</th>
<th>Secondary endpoints</th>
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<tbody>
<tr>
<td>Population included in step 3</td>
<td>PFS will be measured from the time of cross-over to the time of tumour progression (as assessed by the investigator per RECIST V.1.1) or death (whichever comes first)—in all patients undergoing the crossover</td>
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<tr>
<td>To assess PFS in patients undergoing a cross-over following RECIST tumour progression in arm A, from the start of crossover</td>
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<tr>
<td>Population included in step 2 and 3—arm A and B</td>
<td>Time to strategy failure will be measured from the time of randomisation until palbociclib and endocrine therapy discontinuation or death (whichever comes first)</td>
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<td>To assess whether, in patients with rising ESR1 mutations, the early switch (switch following ctDNA detection) to fulvestrant leads a longer time to strategy failure from initial randomisation, than a late switch (cross-over following RECIST tumour progression).</td>
<td>Chemotherapy-free survival will be measured from the time of randomisation until the date of chemotherapy initiation or death (whichever comes first)—in all randomised patients. Anticancer treatments received after the study treatment discontinuation will be described.</td>
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<tr>
<td>To assess whether, in patients with rising ESR1 mutations, the early switch (switch following ctDNA detection) to fulvestrant leads a longer chemotherapy-free survival from initial randomisation, than a late switch (cross-over following RECIST tumour progression).</td>
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<tr>
<td>All included patients—step 1–3</td>
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<td>To report the efficacy (PFS defined on conventional RECIST criteria) of palbociclib combined with hormone therapy (aromatase inhibitor or fulvestrant), from the date of initial inclusion into the trial.</td>
<td>PFS will be measured from the time of inclusion to the time of tumour progression (as assessed by the investigator per RECIST V.1.1) or death (whichever comes first)—in all included patients including those switched to fulvestrant</td>
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<tr>
<td>To obtain additional safety data in a broad patient population treated with palbociclib and hormone therapy (aromatase inhibitor or fulvestrant) in a general oncology practice context</td>
<td>Description of all extrahaematological grade ≥3 toxicities and SAEs incidence rate in the overall population and each treatment step.</td>
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<tr>
<td>To study the patient’s reported quality of life before and until 2 years of therapy</td>
<td>Quality of life score obtained through self-administered QLQ-C30 questionnaire at baseline, at randomisation, and every two cycles until disease progression (including patients who perform a late switch from arm A to B) or until 2 years after inclusion whatever the step if patient did not undergo disease progression or rising ctDNA before 2 years.</td>
</tr>
<tr>
<td>Overall survival</td>
<td>Overall survival measured from the date of inclusion to that of the patient’s death—in all included patients.</td>
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<tr>
<td>Translational objectives (optional)</td>
<td>Translational endpoints</td>
</tr>
<tr>
<td>To report the quantitative and qualitative analyses of circulating tumour DNA detection before and during therapy, the comparison with archived tumour tissue, clinical/pathological characteristics and efficacy of therapy.</td>
<td>ctDNA detection at different time points</td>
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ctDNA, circulating tumour DNA; PFS, progression-free survival; SAE, serious AE.

until randomisation. After randomisation, palbociclib will be given either in association with standard endocrine therapy (letrozole, anastrozole or exemestane) or in association with an ER antagonist (fulvestrant). In the event of significant treatment-related toxicity, palbociclib administration may need to be adjusted. Depending on the nature of the toxicity observed, dosing adjustment may be required for just one or both study drugs in the combination. In the event of significant treatment-related toxicity, palbociclib dosing may be interrupted or delayed and/or reduced as described below. Dose modifications may occur in three ways: (1) Within a cycle: dosing interruption until adequate recovery and dose reduction, if required, during a given treatment cycle; (2) Between cycles: next cycle administration may be delayed due to persisting toxicity when a new cycle is due to start; (3) In the next cycle: dose reduction may be required in a subsequent cycle based on toxicity experienced in the previous cycle. Dose reduction of palbociclib to 100 mg/day, or if needed, to 75 mg/day is recommended depending on the type and severity of toxicity encountered. The use of other concomitant medication/therapy deemed necessary for the care of the patient is allowed.

All medications and therapies taken by the patients or administered to the patients at the onset of trial and all medication given in addition to the IMP during the trial are considered as concomitant medications. The use of other concomitant medication/therapy deemed
Box 1 Complete list of inclusion and exclusion criteria

**Inclusion criteria (STEP1)**

1. Women with proven locoregionally recurrent or metastatic disease adenocarcinoma of the breast not amenable to curative therapy with disease considered potentiely sensitive to aromatase inhibitor. Note: patients relapsing while on adjuvant tamoxifen or other non-aromastase inhibitor adjuvant endocrine therapy and patients relapsing more than 1 year after the end of aromatase inhibitor adjuvant therapy are eligible for this study.
2. Age ≥18 years.
3. Life expectancy >3 months.
4. Eastern Cooperative Oncology Group (ECOG) performance status (PS) 0–2.
5. Oestrogen receptor (ER)-positive and HER2-negative breast cancer. Where available, assessment of ER status should be based on the most recent tumour sample; to be considered as ER-positive, the most recent breast cancer tissue examined must display at least 10% of cancer cells with positive ER staining.
6. Tumour block (primary tumour or metastasis) available.
7. No prior systemic anticancer therapy for metastatic or advanced disease (chemotherapy targeted therapy or endocrine therapy); prior initiation of luteinising hormone-releasing hormone (LHRH) agonist or bone-directed agents is however allowed.
8. Menopausal patients or patients with suppressed ovarian function:
   - Women with bilateral oophorectomy.
   - Postmenopausal women, as defined by any of the following criteria:
     - Age 60 or over.
     - Age 50–59 years and meets one of the following criteria:
       - Amenorrhoea for ≥24 months and follicle-stimulating hormone within the postmenopausal range.
       - Patients with hysterectomy or chemotherapy-induced amenorrhoea must display follicle-stimulating hormone within the postmenopausal range.
   - Other women, provided they are being treated with monthly LHRH analogues (first injection performed ≥7 days before the treatment initiation) and are willing to continue to receive LHRH agonist therapy for the duration of the trial.
9. Patients may have measurable (according to Response Evaluation Criterion in Solid Tumours (RECIST V.1.1) or not measurable disease:
   - Patients with only blastic bone lesions are not eligible.
   - Patients with only pleural, cardiac or peritoneal effusion or measurable nodes with disease considered potentially sensitive to aromatase inhibitor adjuvant endocrine therapy and patients that may require major surgery during the course of the study; however, surgical diagnostic procedure is allowed (even if under general anaesthesia).
   - Patients unable to swallow tablets.
10. Adequate organ and marrow function as defined below:
    - Hqemoglobin ≥90 g/L.
    - Absolute neutrophil count ≥1.5 G/L.
    - Platelet count ≥100 G/L.
    - Serum bilirubin ≤1.5 × ULN (Upper Limit of Normal value). This will not apply to patients with confirmed Gilbert’s syndrome.
    - ALT (alanine aminotransferase) and AST (aspartate aminotransferase) ≤3 × ULN.
    - Alkaline phosphatase ≤2.5 × ULN.
    - Serum creatinine ≤1.5 × ULN or calculated creatinine clearance ≥60 mL/min as determined by Cockcroft-Gault (using actual body weight) formula for females [creatinine clearance=weight (kg) × (140 – age)×0.85 (mL/min)/ (72×serum creatinine (mg/dL))].
11. Willingness and ability to comply with scheduled visits, treatment plan, laboratory tests, and any protocol-related procedures including screening evaluations.

Box 1 Continued

12. Resolution of all acute toxic effects or prior anticancer therapy or surgical procedures to NCI-Common Terminology Criteria for Adverse Events V.4.03 grade 1 (except alopecia or other toxicities not considered a safety risk for the patient at investigator’s discretion).
13. Written informed consent obtained prior to performing any protocol-related procedures including screening evaluations.
14. Patient affiliated to a social security system.

**Non-inclusion criteria (step 1)**

1. Locally advanced breast cancer or locoregional relapse amenable for any treatment with curative intent.
2. HER2-positive or equivocal tumour status either on the primary or on the recurrent tumour, defined as IHC3+, Fish/Cish amplified or Fish/Cish equivocal according to the ASCO2015 criteria.
3. Prior endocrine therapy in the metastatic setting is not allowed.
4. Prior treatment with any CDK 4/6 inhibitor in the adjuvant or metastatic setting (neoadjuvant/preoperative treatment is allowed); however, prior therapy with another targeted treatment in the adjuvant setting is allowed.
5. Visceral crisis: Advanced, symptomatic, visceral spread that is at risk of life-threatening complication in the short term and that requires chemotherapy.
6. Any major surgery (defined as requiring general anaesthesia) or significant traumatic injury within 4 weeks of treatment initiation or patients that may require major surgery during the course of the study; however, surgical diagnostic procedure is allowed (even if under general anaesthesia).
7. Known active, bleeding diathesis.
8. Any serious known concomitant systemic disorder incompatible with the study (at the discretion of investigator), previous history of bleeding diathesis, or anticoagulation treatment (the use of low molecular weight heparin is allowed).
9. Patients unable to swallow tablets.
10. History of mal-absorption syndrome or other condition that would interfere with enteral absorption.
11. Chronic daily treatment with corticosteroids with a dose of ≥10 mg/day methylprednisolone equivalent (excluding inhaled steroids).
12. Known active uncontrolled or symptomatic CNS metastases, carcinomatous meningitis, or leptomeningeal disease as indicated by clinical symptoms, cerebral oedema and/or progressive growth. Patients with a history of CNS metastases or cord compression are eligible if they have been definitively treated with local therapy (eg, radiotherapy, stereotactic surgery) and are clinically stable and off anticonvulsants and steroids for at least 4 weeks before treatment start.
13. Known hypersensitivity to letrozole, anastrozole, exemestane, fulvestrant, palbociclib or any of their excipients.
14. Uncontrolled electrolyte disorders that can compound the effects of a QTc prolonging drug (eg, hypocalcaemia, hypokalaemia, hypomagnesaemia).
15. Patients treated within the last 7 days prior to treatment start in the trial with drug that are known to be CYP3A4 inhibitors, drugs that are known to be CYP3A4 inducers, or with patients who underwent a grapefruit and grapefruit juice cure.
16. Patients already included in another therapeutic trial evaluating an investigational medicinal product or having received an investigational medicinal product within 3 months.
17. History of previous: Continued
2. Patients who have been included in the PADA-1 study, who were randomised for the no-change arm (arm A) on rising ESR1 circulating tumour DNA.
3. Patients who have recent documented tumour progression (RECIST V.1.1).

Non-Inclusion criteria, randomised part (step 2):
1. Patients who have stopped the aromatase inhibitor therapy for more than four consecutive weeks.
2. Patients with a visceral crisis linked to their underlying breast cancer.

Inclusion criteria, crossover part (STEP 3):
1. Patients who have recent documented tumour progression (RECIST V.1.1).

Search for circulating ESR1 mutations
Patients will be access by molecular testing in two regional molecular cancer genetics platforms certified by UNICANCER for the PADA-1 trial.

Mandatory blood samples are required during this study for ESR1 mutations screening in ctDNA. A non-compliance of collection of blood sample for ctDNA measurement will lead to patient withdrawal. For the purpose of the primary objective, blood samples will be collected at inclusion (baseline and/or cycle 1 day 1), after the first cycle (cycle 2 day 1), after the third cycle (cycle 4 day 1) and then every other two cycles from enrolment in the study until the end of treatment. At each blood collection, 20 mL of blood will be drawn into 2 Streck tubes. When possible, blood collections will be performed during routine blood tests. Streck blood tubes will be sent by express carrier to the central laboratory at room temperature within 36 hours and should be received within 96 hours of collection. Blood tubes will be first centrifuged at 1500 g for 10 min at room temperature to pellet cells. The supernatant (‘low-speed’ plasma) will then be harvested (and may be stored at −80°C). In order to extract plasma DNA, ‘low-speed’ plasma will be centrifuged at 16,000 g for 10 min at room temperature to remove cell debris. The supernatant (‘high-speed’ plasma) may be stored at −80°C. Cell-free circulating DNA will be extracted from 4 mL of ‘high-speed’ plasma using the QIAamp circulating nucleic acid kit (Qiagen) according to the manufacturer’s protocol. The extraction procedure can be performed either manually or using automated devices approved by the study biologists. Cell-free circulating DNA may be stored at −20°C. Quantification of ESR1 wild type and mutant (ctDNA) alleles will be performed using ddPCR, an amplification-based technique that detect absolute number of molecules. The PCR reaction will be performed using custom-made or commercially available Taqman assays targeting ESR1 exon 5 and 8 mutations, using the Bio-Rad System. The specificity of each primer pair will be determined on ESR1 DNA in order to define the threshold to discriminate between false and true positive mutant droplets. Each test will contain at least two negative control wells with no DNA and ESR1 DNA as well as an external positive control at 0.5% ESR1 mutant allele frequency (MAF), which will determine the readability of the test. Results will be expressed by the laboratory as the number of ESR1 mutant copies detected per 4 mL of plasma together with the MAF. When the number of mutant ESR1 droplets is null or below the specificity threshold of the assay, the result will be notified ‘no detectable ESR1 mutation’ to the study sponsor. The turn-around time from sample reception by the laboratory to the results should not exceed a median of 10 working days and cannot exceed 15 working days.

Definition of rising ctDNA: At baseline and/or C1D1 (inclusion in step 1) and first visit (after 1 cycle of treatment for example, C2D1, no patient can be considered displaying a rising ctDNA ESR1 mutation. At the following visits (after C4D1 and then every other two cycles), any detectable ESR1 mutation (whatever its level) in patients who had no ESR1 mutation detectable in the previous two blood sampling will be considered as ‘rising ctDNA’. An algorithm is predefined by platform geneticist to decide on the ‘rising’ or ‘non-rising’ status (online supplemental table 1). In patients with detectable levels of ctDNA ESR1 mutation in at least one of the two most recent analyses, ‘rising ctDNA’ detection will be defined for each case by the study biologists, considering levels observed at previous time points and the extent of increase. These cases will be discussed by a board including expert oncologists, biologists (representatives of the central genomic platform) and the principal investigator. Note that these
cases correspond to patients harbouring detectable ESR1 mutation from the very beginning of the study. The number of such patients should be limited, as ESR1 mutation are very uncommon in patients whose cancer is not resistant to AIs.

For translational studies and pending the patient approval (specific consent), all participating patients will be asked to give an additional 20mL of blood (drawn on Streck tubes) for further circulating/ctDNA biomarker research during mandatory ESR1/mutation blood assessments at step 1 (baseline, cycle 2 day 1, cycle 4 day 1, and progression (if applicable)), step 2 (randomisation, cycle 3 day 1, cycle 5 day 1, and progression) and step 3 (cross-over, cycle 3 day 1, cycle 5 day 1, and progression). Furthermore, all patients entering the main study will be proposed to participate to the translational tumour sampling study, whose aim to search for genetic changes driving the early or late resistance to the combination of palbociclib and endocrine therapy. This study will be conducted on archived tumour tissue (inclusion) and archived tumour sample (at progression).

**Visits and follow-up**

Patients will be monitored from the date of inclusion to the date of disease progression or, for patients randomised in arm A and who cross-over after a first disease progression, until the date of disease progression under fulvestrant +palbociclib (second disease progression). Table 2 summarises the follow-up examination/visit schedule. The study is composed of three steps: (1) Step 1 is from the enrolment of the patient (inclusion visit) until the detection of rising ESR1 mutation, or disease progression; (2) Step 2 is from the randomisation until disease progression and (3) Step 3 is a cross-over (only for patients randomised in the arm A and who progressed). This cross-over offers patients initially under palbociclib +AI the possibility to receive palbociclib +fulvestrant.

Eligible patients with signed informed consent form will have an initial assessment at inclusion including clinical evaluation, disease assessment (preferably by CT scan or MRI), standard blood tests, serum biological markers (CA15.3 and CEA) test, ESR1 mutations in ctDNA detection, quality of life (QoL) questionnaires. Physical examination, biological tests including the two serum markers (CA15.3 and CEA) test, ESR1 mutation detection, QoL, and safety assessment will be repeated on the first day of cycle 2 and then every two cycles. At disease progression, patients not randomised or randomised in arm B will stop the study treatment and will be followed during 2 years for their survival status. In Arm A, if considered relevant by the physician, patient will be given the opportunity to perform an optional cross-over to receive palbociclib +fulvestrant until further disease progression. An end of study treatment will occur 30 days after the last day of treatment intake. Following disease progression, the survival status (and the date of death if applicable) and new lines of treatment will be collected every 6 months for 2 years.

**Sample size and power calculation**

The study is a randomised, open label, controlled, phase III clinical trial with two coprimary objectives of safety and efficacy.

To assess whether a change of endocrine therapy associated with palbociclib (namely, an early switch from AI+palbociclib to fulvestrant +palbociclib) will extend the PFS of patients in which rising ESR1 mutations are detected in ctDNA during treatment with palbociclib and AI, we hypothesised: (1) the median PFS of patients whose treatment is unchanged (AI+palbociclib, arm A) will be of 4 months (H0), (2) patients early switching to fulvestrant and palbociclib (Arm B), will display an HR of 0.6 (median PFS of 6.7 months). To highlight such a difference with a power of 80% and a two-sided alpha risk at 5%, 180 patients with rising ESR1 mutation must be randomised to observe 120 events (tumour progression). Because of study drop-offs and eventual technical problems, we anticipate including up to 10% more patients, thus up to 200 patients should be randomised. Based on current scientific reports, we estimate that ESR1 mutations can be detected in the ctDNA of 20% of patients during AI-based therapy. A population of 1000 patients treated by AI and palbociclib is therefore needed to observe up to 200 patients with rising ESR1. The randomisation will be stratified by site of disease (visceral vs non-visceral) and the time to rising ESR1 mutation, defined as the time elapsed between inclusion in the study and the detection of rising ESR1 mutations (≥to 12 months vs <to 12 months).

To obtain additional safety data in a broad patient population treated with palbociclib +endocrine therapy (AI or fulvestrant) and in a general oncology practice context may help to better understand risk factors for palbociclib-related complications. According to the previous PALOMA-2 study (and confirmed by the MONALEESA-2 study with another CDK4/6 inhibitor8), 10% of breast cancer patients presented a one or more episodes of grade 4 neutropenia during therapy with CDK4/6 inhibitors (before tumour progression and treatment discontinuation). If patients with bone metastasis at baseline are exhibiting a 15% rate of grade 4 neutropenia at any time during therapy, this would highlight that this subgroup deserves a particular consideration and/or surveillance. Assuming that advanced breast cancer patients presenting bone metastases is around 60%, 600 patients with bone metastasis at baseline (among 1000 metastases breast cancer patients) will allow to highlight 15% of neutropenia with a precision of 3% (95% CI of ±3%).

Taken together, with the above-mentioned population of 1000 patients, the study will be able to assess both the efficacy of a change of the endocrine therapy associated with palbociclib with 80% statistical power, and to report rate of grade 4 neutropenia with a precision of 3%. The two coprimary objectives will be analysed independently.
### Table 2  Schedule of activity

<table>
<thead>
<tr>
<th>Visits</th>
<th>D0</th>
<th>Cycle 1</th>
<th>Cycle 2</th>
<th>Every two cycles (C4, C6, etc)</th>
<th>Randomisation</th>
<th>Every two cycles (C3, C5, etc)</th>
<th>Cross-over</th>
<th>Every two cycles (C3, C5, etc)</th>
<th>At disease progression</th>
<th>End of study treatment*</th>
<th>Follow-up during 2 years†</th>
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<td>Imaging</td>
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<td>X¶¶</td>
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<td>X d1+d15</td>
<td>X d1+d15</td>
<td>X</td>
<td>XSS</td>
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<td>X</td>
<td>XSS</td>
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<td>X</td>
<td>X</td>
<td>XSS</td>
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<td>CA15-3, CEA</td>
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<td>X (the first 6 months)</td>
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<td>ESR1mut detection (20mL blood)†††</td>
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<td>X</td>
<td>X</td>
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<tr>
<td>Endocrine therapy dispensation</td>
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<td>Fulv. (arm A)</td>
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</tbody>
</table>

**Table continued**
<table>
<thead>
<tr>
<th>Visits</th>
<th>Step 1</th>
<th>Step 2 (at rising ESR1&lt;sub&gt;1m&lt;/sub&gt;)</th>
<th>Step 3 (cross-over)</th>
<th>End of study treatment*</th>
<th>Follow-up during 2 years†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Baseline</td>
<td>Al + palbociclib</td>
<td>Randomisation</td>
<td>Fulvestrant+palbociclib only for pts in arm A</td>
<td></td>
</tr>
<tr>
<td></td>
<td>D0</td>
<td>Cycle 1</td>
<td>Cycle 2</td>
<td>Every two cycles (C3, C5, etc)</td>
<td>At disease progression</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Every two cycles (C4, C6, etc)</td>
<td></td>
<td>Every two cycles (C3, C5, etc)</td>
<td>Every 6 months (±1 month)</td>
</tr>
</tbody>
</table>

Ionogram: sodium, potassium, calcium, magnesium.
Liver and renal functions: creatinine, total bilirubin, lactate dehydrogenase, AST, ALT, ALP, γGT.
Vital signs: heart rate, blood pressure and temperature.
Hemostasis: PT (prothrombin time), PTT (partial thromboplastin time), INR (International Normalized Ratio)

*An end of treatment visit (30 days after the end of study treatment) is required to assess the safety and toxicity of the treatment.
†The post-treatment follow-up will be done every 6 months during 2 years to record information on patient's survival and new line of treatment. Phone calls are allowed for this follow-up period.
‡The patient agrees to the possibility to cross-over in the informed consent at randomisation.
§Results of standard of care tests or examinations performed prior to obtaining informed consent and within 28 days prior to study start (baseline) may be used (for bone scans that period is up to 60 days prior to the start of study treatment).
¶In step 1, tumour evaluation frequency is according to the investigator's choice, with a maximum of four cycles between two tumor assessments.
§§It is not necessary to perform again the tumour evaluation, if the exams were performed less than 28 days prior to the randomisation.
¶¶In steps 2 and 3, tumour evaluation frequency will be performed every two cycles during the first 6 months. After 6 months of the randomisation and until disease progression, the tumour evaluation frequency could be modified according to the investigator choice, with a maximum of four cycles between two tumour assessments.
††It is not necessary to perform again the biological tests if the exams were performed less than 7 days prior to the randomisation or prior to disease progression.
‡‡It is not necessary to perform again the biological tests if the exams were performed less than 7 days prior to the randomisation or prior to disease progression.
***QLQ-C30: completion until 2 years after inclusion if patient did not undergo disease progression or rising ctDNA before 2 years.
****First samples must be drawn at baseline and/or cycle 1 day one prior to the first dose of study drugs. If the sample was performed at baseline, it is not necessary to repeat the samples at C1D1 if the delay between baseline and C1D1 is inferior to 14 days.
†††Fulvestrant or fulvestrant (fulv.) will be dispensed as usual practice.
AI, aromatase inhibitor.
STATISTICAL ANALYSIS

Coprimary outcome
We propose to test superiority in efficacy analysis using two analysis sets; the ‘intention to treat’ (ITT) set, considering all patients randomised regardless of whether they received the randomised treatment, and the ‘per protocol’ analysis set, considering all randomised patients who followed the protocol treatment strategy. The superiority of interventional therapy will be declared only if shown to be superior in the ITT analysis set. The distribution of PFS will be estimated for each intervention arm using Kaplan-Meier method and compared using the log-rank test. A cox model stratified by site of disease (visceral vs non-visceral) and the time to rising ESR1 mutation will be performed to estimate the treatment HR and its 95% CI.

Haematological toxicities of grade ≥3 will be summarised by number and percentage with 95% CI calculated using exact method. The association between incidence of grade ≥3 haematological toxicities and baseline characteristics (Eastern Cooperative Oncology Group Performance Status, metastases localisation, number of metastases site) will be evaluated with χ² test or Fisher’s exact test.

Secondary outcomes
PFS in all screened patients and in patients proceeded in step 3, time to strategy failure and chemotherapy-free survival will be conducted by the Kaplan-Meier method; HR with two-sided 95% CIs will be reported. All extra-haematological grade ≥3 toxicities and serious AEs will be summarised by number and percentage in the overall population and each treatment step. The safety profile of palbociclib +endocrine therapy will be analysed considering all patients that received palbociclib. The EORTC-QLQ-C30 questionnaire will be used to report health-related QoL in all included patients. Summary statistics of the scores for all functional/symptom scales will be calculated at each assessment time point.

Translational outcomes
Number of patients presenting mutations in ctDNA (detection/absence of detection) will be reported with the percentages at each time point. Number of mutant copies and allelic frequencies will be summarised with means, SD, median and range at each time point. Association between ctDNA mutation and clinical factors will be evaluated using χ² test or Fisher exact test, if appropriate, (for ctDNA mutation as qualitative variable), and Student test or Wilcoxon rank-sum test depending on normality of the data (for ctDNA mutation as continuous variable).

DATA MANAGEMENT

The management and the monitoring of the data will be realised according to unicancer procedures and is documented in the monitoring plan. Patients will be considered lost to follow-up if no contact can be established despite repeat attempts to contact. Investigators must document all attempts to contact missing patients throughout the study period unless consent for follow-up has been withdrawn. Furthermore, we will report reasons for withdrawal for each randomisation group and compare the reasons qualitatively. Then we expect to have no missing data for the variables required to construct primary and secondary judgement criteria.

DISCUSSION

Previous research showed that acquired ESR1 mutations in ctDNA are frequently observed after AI exposure in metastatic breast cancer which provide mechanism of resistance to AI therapy. Furthermore, several studies reported the association of ESR1 mutations with poorer PFS and OS. However, no clinical trial has demonstrated the clinical utility of changing treatment at the onset of the ESR1 mutation by dynamically monitoring ctDNA so far. PADA-1 has been designed to test the clinical utility of an early change of treatment after the rise of ctDNA ESR1 mutations in metastatic breast cancer setting. We expect to detect rising ESR1 mutations in patients treated by AI and palbociclib a few months before the actual radiological tumour progression. A coprimary objective of the study is to test whether an early switch from AI+palbociclibto fulvestrant +palbociblib would increase PFS in patients with rising ctDNA ESR1 mutation without radiological progression. Note that only a fraction of the included patient population will be randomised, as (1) ESR1 mutation is not the only mechanism of resistance and (2) for some patient, ESR1 mutational load may be under the limit of detection of the ctDNA-based screening technology used (we considered that 20% of patients will have ESR1 mutation detected in ctDNA).

From a methodological point of view, a fair evaluation of the ‘early change’ of therapy (early switch from AI to fulvestrant before progression), requires a comparison with ‘late change’ (ie, change after tumour progression). To answer this question, we propose a cross-over for patients with ESR1 mutation detection before tumour progression and randomised to the ‘no change’ Arm. Assessing PFS under palbociclib plus fulvestrant after tumour progression on palbociclib plus AI will further improved evidence of the clinical utility of ESR1 mutation detection in real time, as performed in the PADA-1 trial.

The second coprimary objective is the global safety of the combination endocrine therapy +palbociblibl. While efficacy has been demonstrated, the number of patients in which the palbociclib toxicity profile has been deeply assessed remains limited. Palbociclib is considered as a drug with limited toxicity, yet 10% of patients experience grade IV neutropenia during treatment. Our study aims at reporting and analysing treatment toxicity with a focus on the only frequent grade IV toxicity, neutropenia (10% in PALOMA-2 and MONALEESA-2), reported so far with CDK4/6 inhibitors and to analyse this toxicity in regards to the patient baseline characteristics (bone and bone marrow metastasis being eventually involved in that...
toxicity). We preferred to use a precision approach to estimate these toxicity rates rather than a statistical test to avoid spending alpha risk on this endpoint, which would have resulted in a significant increase in the number of patients to be included in this trial aiming to validate an innovative therapeutic strategy based on ctDNA and not directly on the drug itself.

Protocol version

The protocol version number is V.5.0. Submission history is listed in table 3.

Trial status

The first patient was enrolled on 22 March 2017. As of 1 July 2021, 1067 patients have registered in step #1 of the study, 1017 have been included and 172 patients have been randomised in step #2.

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Contributors

FB as principal investigator, SD, TB and A-CH-B conceived and designed the study and helped to draft the manuscript. FB contributed to the statistical design of the study, drafted the manuscript and will analyse the clinical data. SD, A-CH-B, TB, IB, AP, TDLMR, FC, J-LC, FA, LA, JL, MM and SM reviewed this manuscript.

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Disclaimer

Pfizer will not be involved in the collection, analysis, interpretation of the data and writing of scientific manuscripts. Pfizer will have the opportunity to prospectively review any proposed publication, abstract or other type of reports but may not change the conclusions and content.

Competing interests

SD: reports grants and non-financial support from Pfizer, grants from Novartis, grants and non-financial support from AstraZeneca, grants and non-financial support from Roche Genentech, grants from Lilly, grants from Puma, grants from Myriad, grants from Orion, grants from Amgen, grants from Sanofi, grants from Genomic Health, grants from GE, grants from Servier, grants from MSD, grants from BMS, grants from Pfizer, grants from Pierre Fabre, outside the submitted work: A-CH-B: reports personal fees from AstraZeneca, personal fees from Daiichi, personal fees from Clovis, personal fees from GSK, personal fees from MSD, personal fees from Novartis, personal fees from Pfizer, personal fees from Roche outside the submitted work: TB: reports personal fees and non-financial support from Roche, grants, personal fees and non-financial support from Novartis, grants, personal fees and non-financial support from AstraZeneca, grants, personal fees

Table 3: Submission history

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and non-financial support from Pfizer, personal fees from Seagen, outside the submitted work. TDLMR: reports grants, personal fees and non-financial support from Pfizer, grants and non-financial support from Novartis, personal fees and non-financial support from AstraZeneca, personal fees and non-financial support from Roche Genentech, grants and non-financial support from MSD, personal fees and non-financial support from TESARO-GSK, personal fees from CLOVIS ONCOLOGY, personal fees from MYLAN, outside the submitted work. FA: reports grants from Roche, grants from AstraZeneca, grants from Daiichi Sankyo, grants from Pfizer, grants from Novartis, grants from Lilly, outside the submitted work. LA: report personal fees from Roche, personal fees from MSD, personal fees from AstraZeneca, personal fees from BMS, outside the submitted work. FC: reports grants from AstraZeneca, grants, personal fees and non-financial support from Roche, personal fees from Lilly, personal fees and non-financial support from Merck Serono, personal fees and non-financial support from BMS, outside the submitted work. F-CB: reports grants from PFIZER, during the conduct of the study; grants, personal fees and non-financial support from PFIZER, grants, personal fees and non-financial support from NOVARTIS, personal fees from Lilly, personal fees and non-financial support from ROCHE, personal fees and non-financial support from AMGEN, personal fees from SANOFI, personal fees from Radius, grants and personal fees from Seagen, grant from Prolynx outside the submitted work; in addition, F-CB has a patent cDNA detection by ddPCR pending.

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