Serial circulating tumor DNA monitoring of CDK4/6 inhibitors response in metastatic breast cancer

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Abstract
Cyclin-dependent kinase 4/6 inhibitors (CDK4/6i) significantly improve progression-free survival and have become the standard therapy for estrogen receptor-positive/human epidermal growth factor receptor 2-negative metastatic breast cancer patients. Treatment surveillance by radiological imaging has some limitations in detection and repeated biopsy genomic profiling is not clinically feasible. Serial circulating tumor DNA (ctDNA) analysis may provide insights into treatment response. Here we performed serial ctDNA analysis (n = 178) on 33 patients. Serial ctDNA analysis identified disease progression with sensitivity of 75% and specificity of 92%. In eight of 12 patients (61%) responding to CDK4/6i who eventually developed progressive disease, serial sampling every 3 or 6 months captured the initial rise of ctDNA with an average lead time of 3 months. In three of eight patients that did not respond to CDK4/6i (progressive disease at first radiological assessment, 3 months), biweekly sequencing within the first cycle of CDK4/6i treatment (1 month) detected sustained ctDNA levels (≥0.2% variant allele frequency), with lead time of 2 months. Serial ctDNA analysis tracked RECIST response, including clinically challenging scenarios (bone metastases or small-sized target lesions), as well as detecting acquired genetic alterations linked to CDK4/6i resistance in the G1 to S transition phase. Circulating tumor DNA analysis was more sensitive than carcinoembryonic antigen or cancer antigen 15-3 serum tumor markers at monitoring tumor response to CDK4/6i treatment. Our findings indicated the possible clinical utility of serial ctDNA analysis for earlier progressive disease detection and real-time monitoring of CDK4/6i response.

Abbreviations: CA15-3, Cancer antigen 15-3; CDK4/6i, Cyclin dependent kinase 4/6 inhibitor; CEA, Carcinoembryonic antigen; cfDNA, Cell-free deoxyribonucleic acid; cTNA, Cell-free total nucleic acid; CNV, Copy number variant; CR, Complete response; ctDNA, Circulating tumor DNA; dPCR, Droplet digital polymerase chain reaction; dUTP, Deoxyuridine triphosphate; EDTA-2Na, Ethylenediaminetetraacetic acid disodium; ER+, Estrogen receptor positive; FDA, U.S. Food and Drug Administration; HER2-, Human epidermal growth factor receptor 2 negative; HR, Hazards ratio; IRB, Institutional review board; MBC, Metastatic breast cancer; NGS, Next generation sequencing; NPA, Negative percent agreement; PCR, Polymerase chain reaction; PD, Progressive disease; PFS, Progression free survival; PR, Partial response; RECIST, Response Evaluation Criteria in Solid Tumors; SD, Stable disease; SNV, Single nucleotide variant; TMAP, Torrent mapping alignment program; UMT, Unique molecular tags; VAF, Variant allele frequency; WBC, White blood cell.
1 | INTRODUCTION

Cyclin-dependent kinase 4/6 inhibitors have become the standard treatment for ER+/HER2− advanced breast cancer patients. Preclinical studies have established the role of a CDK4/6–RB pathway in regulating the cell cycle through the G1 to S transition phase.1,2 Multiple clinical trials have reported that patients receiving CDK4/6i showed significantly longer PFS in combination with endocrine therapy in ER+ breast cancer.3,4 Despite improvements in clinical management, a subset of breast cancer patients will develop resistance to treatment.

Treatment response is mainly assessed by serial radiographic images.5 Radiographic imaging provides assessment of the disease burden but can sometimes be hampered by suboptimal detection limit and inconsistencies in tumor size measurements.6,7 Carcinoembryonic antigen and CA 15-3 are serum biomarkers mainly used for treatment monitoring in patients with MBC,8 with elevated levels often associated with poor prognosis. Although clinically useful in some breast cancer patients, CEA and CA 15-3 have limited sensitivity of only 50%9 and 60%–70%,8,10 respectively. Tumor genome sequencing remains the gold standard for genomic profiling, providing insights into potential actionable targets for personalized treatment.11 However, in a metastatic setting, repeated biopsy is not clinically feasible. Taken together, a more sensitive and less invasive approach is needed to provide an earlier measure of treatment response to inform individual treatment decisions.

Many cancers shed DNA into the bloodstream, referred to as ctDNA. Liquid biopsy through serial ctDNA analysis can complement radiographic imaging and CEA and CA15-3 markers in assessing CDK4/6i responses. It is minimally invasive, feasible for repeated sampling, and is able to detect tumor-specific alterations in circulation, reflecting disease burden in real time.12 Previous studies have investigated the clinical utility of liquid biopsy in predicting response to CDK4/6i.13-15 However, these studies were focused on ctDNA dynamics at early time points14,15 or on the heterogeneity of the mutational landscapes between baseline and at disease progression.15 None of the studies used monitoring through a granular time series to catch disease progression at an earlier time point.

In this single center, retrospective, exploratory biomarker study, we evaluated serial ctDNA analysis against radiographic imaging and CEA and CA 15-3 markers as a biomarker for CDK4/6i response monitoring. We show that short interval serial ctDNA analysis was able to catch initial rise or reemergence of ctDNA, indicative of disease progression, well in advance of radiological assessment. This study also identified mutations potentially predictive of CDK4/6i response, tracked tumor response of RECIST-defined nonmeasurable target lesions, and identified acquired genetic alterations linked to CDK4/6i resistance. Our results support the possible clinical utility of serial ctDNA analysis for earlier PD detection and real-time monitoring of CDK4/6i response.

2 | MATERIALS AND METHODS

2.1 | Patient cohort

Thirty-three ER+/HER2− female MBC patients scheduled to receive CDK4/6i and endocrine combination therapy either as a first line or subsequent line treatment were recruited in a consecutive, retrospective manner from the Cancer Institute Hospital of Japanese Foundation for Cancer Research from April 16, 2018 to September 30, 2020. Patient age varied from 35 to 73 years (median, 54 years) and were eligible if they had progressed on endocrine treatment for metastatic disease or during adjuvant therapy. Study has been approved by the ethical committee of Japanese Foundation for Cancer Research (Institutional Review Board No. 2015-1056), with approval granted for collection and genomic profiling of patient samples. All 33 patients in the study provided written informed consent for cell-free DNA and white blood cell sequencing, with the option to be excluded from the study at any time. No patient withdrew consent for the study. The majority of patients (30/33, 91%) received oral palbociclib 125 mg daily for 3 weeks on, and 1 week off protocol in combination with either fulvestrant (500 mg every 4 weeks), letrozole (2.5 mg daily), tamoxifen (20 mg daily), anastrazole (1 mg daily), or exemestane (25 mg daily). The remaining three patients received oral abemaciclib 300 mg daily in combination with fulvestrant (500 mg every 4 weeks) or anastrazole (1 mg daily). Premenopausal women additionally received goserelin.

Patient records were reviewed to obtain radiological images of computed tomography scans as well as tumor size measurements for the patients receiving CDK4/6i therapy in this study. Radiologic images were obtained at pretreatment and at all subsequent intervals until clinical progression, 24 months evaluation, or the most recent assessment by September 30, 2020 (Figure 1). Radiologic images of multiple target lesions were obtained when available and evaluated at every subsequent interval. Tumor response was assessed and defined by the study clinician according to RECIST version 1.1.5 Patients who developed clinical progression at first radiographic assessment were defined as not responding to CDK4/6i and endocrine combination therapy. Tumor marker levels of CEA and CA 15-3 were obtained retrospectively from patient records at concurrent or nearest to the liquid biopsy sampling time points for comparison.
2.2 Sample processing, library preparation, and sequencing

Blood samples were collected at pretreatment, 2 weeks, and 1, 3, 6, 9, 12, 18, and 24 months or until clinical progression, whichever comes first (Figure 1). For patients with no ctDNA mutations detected at the end of cycle 1 of CDK4/6i therapy (1M), the end of evaluation sample is sequenced to assess for detectable ctDNA mutations. If mutations are present, cfDNA from all sampling intervals are evaluated. Based on our data cut-off on September 30, 2020, we collected 178 plasma samples from 33 patients. Collection and processing of whole blood has been described in previous publications. Briefly, 14 mL whole blood was collected using EDTA-2Na tubes (Terumo). Whole blood was centrifuged at 2000 g at 4°C for 10 minutes to separate plasma from white blood cells and red blood cells. The plasma layer was further centrifuged at 16 000 g at 4°C for 10 minutes to remove cell debris. Both plasma and white blood cells were stored at −80°C until nucleic acid extraction.

Plasma cfTNA was extracted using MagMAX Cell-Free Total Nucleic Acid Isolation kit (Applied Biosystems) according to the manufacturer’s protocol. Genomic DNA from white blood cells was extracted using FlexiGene DNA Kit (Qiagen). Targeted NGS for cfTNA was carried out using Oncomine Pan-Cancer Cell-Free Assay following the manufacturer’s protocol (Ion Torrent). Oncomine Pan-Cancer Cell-Free Assay is an amplicon-based ctDNA targeted assay, incorporating short oligonucleotides called UMTs to distinguish PCR/sequencing artifacts from actual variants (Figure 1). Library construction and subsequent NGS of cfTNA and genomic DNA were undertaken as previously described. In most samples, 20 ng cfTNA was used for library construction. Sequencing of genomic DNA was carried out using the same methods using 30 ng genomic DNA for library construction.

2.3 Analysis of sequencing data

Sequencing reads were aligned to hg19 and variant calling was carried out using Torrent Suite 5.10.1 and Ion Reporter 5.10.3.0 software, respectively. Briefly, raw sequence files were aligned to hg19 using Torrent Mapping Alignment Program (TMAP) with default analysis parameters. Valid NGS runs have 90% or more of total reads mapped, alignment quality with read error rate of 2% or less (AQ17). The targeted minimum total coverage for each amplicon is at least 20,000 X. Library conversion rate was evaluated using the theoretical assumption that 10 ng cfDNA would be equivalent to approximately 3000 haploid genomes. Variant calling was undertaken using the Oncomine TagSeq Pan-Cancer Liquid Biopsy w2.1 workflow with modifications. In order to reduce false positive detection in the plasma ctDNA assay, samples are defined as ctDNA-positive if the following criteria are met: (a) a minimum of three reads with the same UMT is required to form a functional family; (b) SNVs and indels are detected with a minimum of two variants supporting functional families at VAF of 0.2% or more; (c) SNVs and indels are not detected with any counts in the corresponding white blood cells; and (d) minimum of 2.5-fold change...
between CNV target amplicons and non-CN V reference amplicons is required. Variants were annotated using Oncomine Pan-Cancer Annotations version 1, a proprietary list of databases.

2.4 | Digital droplet PCR analysis

For patient CDK28, plasma cfDNA was insufficient for targeted NGS in 1M, 3M, and 6M samples. Circulating tumor DNA mutations were evaluated focusing only on ESR1 Y537S detected by Oncomine Pan-Cancer Cell-Free Assay at disease progression. Digital droplet PCR was carried out for all seven time points for patient CDK28 from pretreatment to disease progression (10.5M). The assay used was LBx Probe Catalog no. A082 (65117) ESR1 multi probe targeting Y537N/S, D538G (Riken Genesis). All reactions were carried out on a QX200 ddPCR system (Bio-Rad). A total of 10 ng cfDNA was used in each PCR reaction consisting of primers, probes, and ddPCR Supermix for probes (no dUTP). Reactions were divided into approximately 20,000 droplets per cell using the QX200 droplet generator. Emulsified PCR was carried out on a 96-well thermal cycler using the following cycling conditions: (1 cycle of 95°C for 10 minutes; 40 cycles of 94°C for 30 seconds, 60°C for 1 minute, one cycle of 98°C for 10 minutes, and 4°C hold). Samples were analyzed using the QuantaSoft software to assess number of droplets harboring mutant ESR1 or WT ESR1.

2.5 | Statistical analysis

The primary outcome was to assess whether granular sampling time series ctDNA analysis could indicate earlier treatment response compared to RECIST-based clinical diagnosis. Secondary outcomes include identifying ctDNA biomarkers predictive of CDK4/6i response, evaluating ctDNA surveillance of clinical responses to CDK4/6i compared to radiological images, CEA and CA15-3 markers, and identifying acquired ctDNA alterations linked to treatment resistance. Circulating tumor DNA monitoring markers are defined as follows: (a) ctDNA detected within cycle 1 of CDK4/6i treatment – within 1 month; (b) ctDNA of 0.2% or more for SNVs and at 2.5-fold or more for CNVs at least once throughout the monitoring period; and (c) ctDNA alteration was not detected with any counts in the corresponding WBC sequencing. Acquired ctDNA alterations are defined as follows: (a) ctDNA undetected within the first cycle of CDK4/6i treatment (SNV count = 0 or CNV = 2.0-fold); or (b) at disease progression, SNV of 0.2% or more or CNV of at least 2.5-fold. For sensitivity and specificity of ctDNA-detected disease progression: (a) ctDNA levels at point of progression shows a minimum of 0.2% for SNVs and at least 2.5-fold for CNVs. Sensitivity is defined as (true positives/(true positives + false negatives)) and expresses the fraction of ctDNA positive patients at point of progression made using plasma targeted NGS. Specificity, or negative percent agreement is defined as (true negatives/(true negatives + false positives)), and expresses the ratio of patients expected to be ctDNA negative at the end of evaluation by plasma targeted NGS. To define lead time for ctDNA analysis, we set the following criteria: (a) SNVs of 0.2% or more or CNVs of 2.5-fold or more in consecutive intervals from initial rise until progressive disease; (b) lead time calculated from 1 month onwards (end of cycle 1 treatment); and (c) lead time is calculated from ctDNA initial rise to nearest sampling interval to clinical progression. Clinical implications of acquired ctDNA alterations were annotated by OncoKB. Association of pretreatment markers with CDK4/6i response was evaluated using Kaplan-Meier and Cox regression analyses, with response time measured from time of CDK4/6i treatment initiation to RECIST-defined clinical progression or tumor response by cut-off date of September 30, 2020. Multivariable Cox regression analysis was used to test the independent prognostic value of pretreatment markers, adjusted for age, presence of visceral metastases, prior lines of endocrine therapy, and prior lines of metastatic therapy. These variables were preselected as known prognostic factors. Kaplan-Meier and Cox regression analyses were undertaken using the “survival” and “survminer” packages in R version 4.0.3. Randomization and blinding were not applicable to our study; as this was a biomarker observational and comparison study, patients were recruited irrespective of the CDK4/6i or combination endocrine therapy they received. Comparison of ctDNA and detection rates of tumor markers CEA and CA 15-3 were evaluated using Fisher’s exact test. All statistical analyses were carried out in R version 4.0.3 using the default statistical package. All statistical tests were two-sided, with P-values less than .05 considered statistically significant.

3 | RESULTS

3.1 | Patient characteristics

By the data cut-off date of September 30, 2020, 178 plasma samples from 33 patients (average, 54 years [35–73 years]) had been recruited with a median follow-up period of 12 months (1–24 months; Table 1). Complete response was observed in one patient and partial response in nine patients, giving an objective response rate of 30% (10 of 33 patients). Sixteen patients had stable disease (Table 1). Eight patients (24%) had no clinical benefit (PD at first radiological assessment) while 12 patients developed disease progression during the follow-up period (Figure 2A). The majority of patients received palbociclib in combination with endocrine therapy (91%; 30/33). At pretreatment, 17 patients (52%) had visceral metastasis, 16 patients (48%) had nonvisceral metastasis with eight of the 16 patients presenting only bone metastasis. All 33 patients were evaluable according to RECIST 1.1 with 13 patients (39%) having measurable disease and 20 (61%) having nonmeasurable disease.

3.2 | Cell-free DNA isolation and ctDNA NGS analysis

Cell-free DNA was successfully extracted from all 178 samples of 33 patients (average, 49.8 ng per 14 mL blood) with 175 of 178 samples
| Description | Clinical and pathological features |
|-------------|----------------------------------|
| Number of patients | 33 |
| Median age, y (range) | 54 (35–73) |
| Pathology (n = 33) | Invasive ductal carcinoma 32 (97), Others 1 (3) |
| Receptor status (n = 33) | ER+PR+ HER2− 30 (91), ER+PR− HER2− 3 (9) |
| Tumor response (n = 33) | Complete response 1 (4), Partial response 9 (27), Stable disease 15 (45), Progressive disease 8 (24) |
| Menopausal status (n = 33) | Premenopausal 12 (36), Postmenopausal 21 (64) |
| CDK4/6i (n = 33) | Palbociclib + endocrine therapy 30 (91), Abemaciclib + endocrine therapy 3 (9) |
| Visceral metastases (n = 33) | Yes 17 (52), No 16 (48) |
| RECIST criteria (n = 33) | Measurable 13 (39), Nonmeasurable 20 (61) |
| Prior lines of endocrine therapy (n = 33) | 0 line 5 (15), 1 line 13 (39), 2 lines 5 (15), 3+ lines 10 (31) |
| Prior lines of metastatic therapy (n = 33) | 0 line 12 (36), 1 line 10 (30), 2 lines 4 (12), 3+ lines 7 (22) |
| ctDNA median follow-up, mo (range) | All patients (n = 33) 12 (1–24), Responders (n = 13) 18 (6–24) |

Note: Data are shown as n (%) unless otherwise noted.
Abbreviations: CDK4/6i, cyclin-dependent kinase 4/6 inhibitor; ctDNA, circulating tumor DNA; ER, estrogen receptor; HER2, human epidermal growth factor receptor 2; PR, progesterone receptor.

Prior lines of endocrine therapy include adjuvant and metastatic regimens prior to CDK4/6i treatment.

Table 1: Clinical characteristics of 33 metastatic breast cancer patients at pretreatment.

Our study adopted a granular time series for early disease progression detection. Serial sampling every 3 or 6 months post cycle 1 of CDK4/6i treatment was able to capture the initial rise of ctDNA earlier than clinical progression. In 8 of 12 patients responding to CDK4/6i (≥6 months) but eventually developing PD, ctDNA analysis detected earlier treatment response, with initial rise or reemergence of ctDNA observed prior to RECIST-defined disease progression (average lead time, 3 months; range, 1.5–6 months; Figures 3 and S2). In three of eight patients that did not respond to CDK4/6i (PD at first radiologic assessment ~3 months), biweekly sequencing within the first cycle of CDK4/6i treatment (1 month) detected sustained ctDNA levels (≥0.2% VAF), with an average lead time of...
2 months (Figures 3 and S2). Based on our findings, ctDNA analysis, through granular sampling intervals, detected tumor-derived mutations indicative of disease progression well in advance of radiologic assessments.

Carcinoembryonic antigen and CA 15-3 are the most widely used serum markers in patients with MBC. From our findings, ctDNA analyses were more sensitive and reflective of tumor response compared to CEA or CA15-3. More patients (20/33 patients, 61%) had detectable ctDNA alterations in at least one time point compared to elevated CEA (>5 ng/mL: 12/33 patients, 36%) or CA15-3 levels (>30 U/ml: 15/33 patients, 45%; Figures S2 and S3), although this trend is suggestive and not statistically significant (P_Fisher > .05).

3.5 Circulating tumor DNA analysis to monitor small-sized tumors, bone metastasis, and heterogenous tumors

Consistent with previous findings, ctDNA analysis concordantly tracked tumor response of RECIST-measurable disease (Figure 4). However, for more challenging scenarios, such as patients showing only bone metastasis or small-sized tumors below RECIST thresholds, the comparisons are less clear. In our study, 11 patients were defined as having nonmeasurable disease at pretreatment due to small tumor sizes that were below RECIST criteria (Table S3). Eight of the 11 patients had detectable ctDNA alterations. In patient CDK05 with tumor sizes below RECIST criteria, ctDNA analysis detected multigene genomic amplifications in FGFR3, FGFR1, MYC, and CCND1 at pretreatment. These CNVs reduced to basal levels during treatment, and reemerging during clinical progression at levels similar to pretreatment (Figure 5A). The ctDNA profiles in patient CDK05 were consistent with presence of liver target lesions observed at both pretreatment and disease progression. Similarly, in patient CDK14, driver mutations PIK3CA H1047R and TP53 Y220C detected at pretreatment increased in tandem with the lung target lesion until clinical progression (Figure 5B). Circulating tumor DNA analysis could be useful considering challenging and potentially inconclusive tumor response interpretation for patients who were not assessable using RECIST criteria.
Monitoring bone metastasis is challenging due to difficulty in interpreting radiologic images or in obtaining tissue biopsies. In the nine patients presenting with only bone metastasis at pretreatment, ctDNA analysis detected breast cancer-related alterations in 67% (6 of 9) of patients (Table S3). In 4 of 6 bone metastasis patients (CDK04, CDK07, CDK16, and CDK21) with detected ctDNA alterations, ctDNA analysis detected rapid increase of pretreatment alterations prior to or at clinical progression, implying detection of possible tumor-specific alterations reflecting resistance towards CDK4/6i treatment (Figure 6). Particularly for patients CDK04, CDK07, and CDK21, increase in pretreatment ctDNA alterations at clinical progression coincided with acquisition of new metastatic lesions in the liver and lung (Figure 6A,B,D). As for patient CDK16 (Figure 6C), pretreatment ctDNA mutation SF3B1 K700E reemerged and has shown rapid increase in ctDNA levels while tumor response continues to be defined as stable disease based on bone metastasis target lesion. Our findings highlight the potential clinical utility of liquid biopsy in complementing radiologic assessment of tumor response when interpretation is challenging or inconclusive.

Circulating tumor DNA analysis has the advantage of capturing genomic heterogeneity of multiple tumor lesions. In patient CDK15, a total of eight ctDNA mutations were detected within cycle 1 of treatment for serial monitoring, comprising of six PIK3CA and two ESR1 mutations (Figure S4). Diverse tumor responses were observed: ctDNA profiles 1, 2, and 4 show constant residual amounts of ctDNA variants throughout CDK4/6i treatment, followed by subsequent increase of ctDNA levels prior to clinical progression. The ctDNA analysis suggests partial inhibition of these tumors, eventually leading to resistance and clinical progression. Circulating tumor DNA profile 5 shows acquisition of AKT1 and PIK3CA mutations, previously undetected at pretreatment, near to or at clinical progression, possibly reflecting acquisition of the new liver lesion at disease progression.

3.6 | Circulating tumor DNA analysis of acquired mutations associated with resistance

In some patients, acquired mutations might emerge due to treatment pressure. Evaluation focused on 12 of 20 patients who initially responded to therapy but later developed disease progression. Serial ctDNA analysis detected seven acquired genetic alterations from five patients in genes from several signaling pathways related to cell cycle checkpoint. Figure 7 shows acquired variants ERBB2 (HER2 tyrosine kinase pathway), ESR1 (estrogen receptor signaling) as well as in downstream genes such as PIK3CA (PI3K-AKT signaling), MYC, and cyclin D1 (cell cycle checkpoint). These genes are known to play critical roles in the G1 to S transition in the cell cycle. We define “acquired variants” as ctDNA that was undetected within first cycle of CDK4/6i treatment (SNV count = 0 or CNV = 2-fold) and detected at disease progression (SNV 0.2% or CNV 2.5-fold). The VAF or copy numbers of paired pretreatment and PD time points are listed in the colored boxes directly below the detected variants. Five of the seven acquired alterations have clinical relevance as potential actionable targets for molecular-based therapeutics, with ERBB2 amplification and PIK3CA N345K being FDA recognized biomarkers predictive of response to the corresponding therapy while AKT1 E17K, ESR1...
DS38G, and ESR1 Y537S show compelling clinical evidence predicting response to the targeted therapy (Table S4). We show some key examples. In patients CDK13 and CDK28, acquired ESR1 mutations, undetected in pretreatment samples, appeared prior to disease progression (Figure S5). For patient CDK13, ESR1 mutation emerged when pretreatment liver target lesion increased in size leading up to disease progression (Figure S5). As for patient CDK28, ESR1 mutation was detected when a new liver lesion, previously undetected at pretreatment, was observed (Figure S5).

4 | DISCUSSION

In this retrospective, longitudinal monitoring study of CDK4/6i therapeutic response, we show serial ctDNA analysis to be a highly informative biomarker that accurately captures the clonal dynamics of MBC tumors. This study presents comprehensive evaluation of MBC response to CDK4/6i through short interval serial ctDNA analysis, using targeted NGS in every interval (175 samples for 33 patients). Both DNA and RNA extraction was carried out as the ctDNA assay can detect somatic alterations as well as gene fusions from plasma. This approach did not impact recovery of cfDNA, achieving an average library conversion rate of 76%.

Our study brings additional insights regarding the biology of CDK4/6i response. Through granular time series ctDNA analysis, we were able to capture the initial rise of ctDNA levels, indicative of disease progression, well in advance of radiological assessment, in patients responding to CDK4/6i treatment. In addition, for patients who did not respond to CDK4/6i, sustained ctDNA detection in consecutive intervals within cycle 1 of CDK4/6i treatment enabled possible earlier detection of disease progression. The potential of ctDNA to detect early disease progression is crucial as it could provide an opportunity for earlier termination of CDK4/6i treatment, avoiding unnecessary CDK4/6i-related side effects. The clinical implications of earlier detection of disease progression remains a contentious point for discussion, with very little data to support claims of improved clinical outcome. However, early data from the PADA-1 study lend support to this claim. PADA-1 is an ongoing ctDNA-based interventional clinical trial, evaluating the efficacy of switching therapy upon detection of plasma ESR1 mutations in ER+HER2- MBC. In the PALOMA-3 study, when evaluating the predictive ability of baseline plasma ESR1 mutations on control arm (fulvestrant and placebo), results show shorter response for mutant ESR1 carriers (ESR1 mutants median PFS, 3.6 months [95% CI, 2.5-5.4 months]; ESR1 WT median PFS, 5.5 months [95% CI, 3.5-7.4 months]; P = .04). In our study, we did not include a control arm (patients receiving only endocrine therapy) to evaluate predictive ability of baseline ESR1 mutations towards treatment efficacy. However, results from the PALOMA-3 trial indicate that patients receiving only endocrine therapy, carrying ESR1 mutations at baseline, will not benefit from treatment, as shown by the shorter median PFS. In our study, patients carrying ESR1 mutations prior to treatment were unlikely to benefit from the CDK4/6i endocrine combination therapy, with the trend remaining statistically significant even after adjusting with known prognostic factors. Although potentially interesting, we acknowledge that the prognostic significance of ESR1 is suggestive and lacks external validation in a larger independent dataset.

Currently, clinical response of solid tumors is assessed by RECIST 1.1 criteria with target lesions selected based on size and suitability for repeated evaluations. Our results imply that the dynamic range of serial ctDNA analysis are concordant with radiologic assessments, supporting the adoption of ctDNA as a monitoring biomarker. When RECIST criteria is not applicable (bone lesions or target lesions below RECIST size), the mutation profiles detected from serial ctDNA analysis could complement radiographic imaging in providing a more decisive assessment of tumor response. In our analysis, we observed that liquid biopsy provided an earlier measure of treatment response, with increasing ctDNA levels detected before radiographic assessment of disease progression or changes in CEA and CA 15-3 levels (Figures S2 and S3). This observation has been supported by other recent studies.
(A) CDK05

Liver lesion
-2 weeks before treatment

New liver lesion emerged
3M - PD

Sternum bone meta
2 weeks before treatment

Sternum bone meta
3M - PD

(B) CDK14

Lung lesion
-1.5M before treatment

Lung lesion
3M - SD

Lung lesion
6M - SD

Lung lesion
10M - PD

Sternum bone meta
-1.5M before treatment

Sternum bone meta
3M - SD

Sternum bone meta
6M - SD

Sternum bone meta
10M - PD
In this study, ctDNA analysis detected acquisition of new alterations in ERBB2, PIK3CA, AKT1, MYC, CCND1, and ESR1. These alterations might have emerged due to prolonged exposure to the CDK4/6i and endocrine combination therapy. These mutated genes are involved in dysregulation of key pathways upstream of the CDK4/6-RB1, possibly rendering cancer cells resistant to CDK4/6 inhibition, in line with previous findings.13,28 ESR1 mutations not only have significant clinical implications at the pretreatment stage, but also at disease progression as a possible mechanism in evading CDK4/6 and endocrine therapy inhibition. Some acquired alterations are clinically relevant as potential actionable targets for molecular based therapeutics, possibly aiding treatment selection after CDK4/6i treatment resistance.

Our study has its limitations. This study was designed to evaluate the clinical validity of liquid biopsy to monitor response to CDK4/6i. A larger population cohort is needed to confidently establish the association of ESR1 mutations towards CDK4/6i response. Further refining the granular time series and a larger cohort will help establish the clinical significance of ctDNA-detected earlier treatment response and the key landmark intervals predictive of CDK4/6i response. The ctDNA alterations detected were largely known breast cancer associated markers. This is due to the ctDNA assay, with targeted alterations largely limited to known mutations in frequently occurring cancers to maximize assay sensitivity. This probably explains the lack of ctDNA detection in some patients, particularly CDK12 and CDK24 who rapidly developed PD. With genes that do...
not harbor hotspot mutations (eg, *RB1*), using hybrid-capture assays covering more target regions at landmark intervals to guide ctDNA monitoring might improve detection sensitivity. Nevertheless, our findings have indicated possible clinical utility of serial ctDNA analysis for early detection of disease progression as well as real-time monitoring of CDK4/6i response. Serial monitoring using targeted NGS liquid biopsy reveal a multifaceted view of MBC clonal evolution during CDK4/6i treatment. The findings from this study are crucial to advancing disease management of advanced breast cancer patients, in particular patients receiving CDK4/6i treatment.

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DISCLOSURE
Y.M.C. reported employment with Cancer Precision Medicine, Inc. Japan outside the submitted work. Y.N. reported receiving advisory fees and stocks from OncoTherapy Science, Inc. Japan outside the submitted work. S.-K.L. reported receiving advisory fees from Cancer Precision Medicine, Inc. Japan outside the submitted work. The remaining authors declare no potential conflicts of interest.

DATA AVAILABILITY STATEMENT
All data supporting the findings of this study are available within the article and its supplementary information/data files, and from the corresponding author upon reasonable request.

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**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of the article at the publisher’s website.

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