Prospective Cohort Study of Combination Therapy With Abemaciclib and Hormonal Therapy for Chemotherapy-Treated Patients With Hormone Receptor-Positive Metastatic Breast Cancer

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Abstract

Background: Combination therapy with cyclin-dependent kinase 4 and 6 (CDK4/6) inhibitors and hormonal therapy as the first-line and second-line treatments has already been shown to be effective in patients with hormone receptor-positive (HR+) human epidermal growth factor receptor 2-negative (HER2-) metastatic breast cancer (MBC) in clinical trials. On the other hand, in clinical practice, CDK4/6 inhibitors are used not only as first-second-line but also as later-line hormonal therapies, or for patients receiving prior chemotherapy in metastatic setting. However, the efficacy and safety of combination therapy in these patients remain unclear. In this study, we evaluate the clinical efficacy and safety of combination therapy with abemaciclib and hormonal therapy for chemotherapy-treated patients with HR+ HER2- MBC.

Methods: This multi-institutional prospective cohort study will involve a total of 300 chemotherapy-treated patients with HR+ HER2-MBC. The primary endpoint is progression-free survival (PFS). Secondary endpoints include overall survival, time to treatment failure, response rate, clinical benefit rate, and adverse events. The preplanned subpopulation analysis is the number of chemotherapy regimens for HR+ HER2- MBC (two or less vs. three or more), prior treatment history with CDK4/6 inhibitors other than abemaciclib (presence vs. absence) and menopausal status (pre vs. post). We also planned to determine PFS of the subpopulation treated with abemaciclib as maintenance therapy after chemotherapy.

Discussion: In this multi-institutional prospective cohort study, we evaluate the clinical efficacy and safety of combination therapy with abemaciclib and hormonal therapy for chemotherapy-treated patients with HR+ HER2- MBC. We also evaluate this combination therapy as maintenance therapy in patients who respond to early-line chemotherapy.

Keywords: Abemaciclib; Hormone receptor-positive metastatic breast cancer; Overall survival; Real-world evidence; Prospective cohort study; Chemotherapy-treated

Introduction

A hormone receptor-positive (HR+) human epidermal growth factor receptor 2-negative (HER2-) subpopulation accounts for two-thirds of metastatic breast cancer (MBC) in Japan [1]. Because of the rarity in achieving cure in these patients, the main purpose of treatment is to palliate symptoms, extend survival, and improve quality of life.

Cyclin-dependent kinase 4 and 6 (CDK4/6) inhibitors are oral molecular-targeted agents that suppress tumor proliferation through specific inhibition of CDK4 and CDK6. In clinical trials, combination therapy with CDK4/6 inhibitors and hormonal therapy as the first-line and second-line treat-
ments is extremely effective in patients with HR+ HER2-MBC. Phase III clinical trials of CDK4/6 inhibitors, such as palbociclib [2, 3], abemaciclib [4, 5], and ribociclib [6-8], have shown significant prolongation of progression-free survival (PFS) using combination therapy compared with hormonal therapy alone. Moreover, abemaciclib and ribociclib in combination with hormonal therapy have significantly prolonged overall survival (OS) [9-11]. The adverse events (AEs) caused by CDK4/6 inhibitors appear to be within the acceptable range. Thus, combination therapy with CDK4/6 inhibitors and hormonal therapy is recommended by the National Comprehensive Cancer Network (NCCN) guidelines [12] and The Japanese Breast Cancer Society Clinical Practice Guideline for Breast Cancer 2018 [13]. In Japan, palbociclib and abemaciclib were respectively listed on the National Health Insurance drug price list in December 2017 and November 2018. Ribociclib was listed as off label in July 2020 in Japan.

However, the advent of CDK4/6 inhibitors has complicated the treatment of HR+ HER2-MBC. Although the effectiveness of CDK4/6 inhibitors appeared to have been confirmed, treatment with a single hormonal agent is still one of the options, and there remains another clinical question about treatment sequence among hormonal agents, molecular targeted agents, and less toxic chemotherapeutic agents. This may be due to the high cost of molecular targeted agents and the uncertainty of their effectiveness when considering the sequential order.

Furthermore, there is a lack of phase 3 studies that verify the efficacy and safety of these agents with hormonal therapy as third-line or later treatments and for patients treated by chemotherapy. As for administration after chemotherapy, palbociclib in combination with second-line or third-line hormonal therapy has been reported in the PALOMA-3 trial [3], and ribociclib has been verified in combination with first-line hormonal therapy in the MONALEESA-7 trial [8]. Abemaciclib has not yet been verified for patients after chemotherapy in phase 3 trials; chemotherapy-treated cases were excluded in the MONARCH-2 trial [4] and the MONARCH-3 trial [5].

Objectives

The objective of this study is to clarify the efficacy and safety of combination therapy with endocrine therapy and abemaciclib in patients who have treatment history with chemotherapeutics for MBC.

Hypotheses

1) Abemaciclib therapy is safe and effective for patients with MBC previously treated with chemotherapeutics.

2) Abemaciclib therapy is an effective maintenance therapy after chemotherapy.

3) Abemaciclib therapy is effective for patients who have been previously treated with other CDK4/6 inhibitors.

Materials and Methods

Study design

This research is a multi-institutional prospective cohort study to evaluate the clinical efficacy and safety of combination therapy with abemaciclib and hormonal therapy for chemotherapy-treated patients with MBC. In addition, we designed three pre-planned analyses evaluating the efficacy and safety of combination therapy with abemaciclib and hormonal therapy for chemotherapy-treated patients with MBC. One is to evaluate maintenance therapy after the chemotherapy which was discontinued for reasons other than disease progression. The other evaluates subpopulations stratified by the number of prior chemotherapy regimens (one or two vs. three or more) and, the third, stratified by prior treatment history with CDK4/6 inhibitors other than abemaciclib. The study design flowchart is shown in Figure 1.

We will also prospectively investigate drug-induced lung injury caused by abemaciclib as an ancillary study.

Study setting

The study was approved by the Protocol Review Committee of the Comprehensive Support Project for Oncological Research of Breast Cancer (CSPOR-BC). This study was registered at the UMIN Clinical Trials Registry as UMIN 000037395. The study was started in December 2019 and will be completed in November 2022. Patient enrollment will be completed in November 2022 (extended 1 year due to insufficient registration).

Endpoints

The primary endpoint is PFS, defined as the time from the date of registration to disease progression or death from any cause, whichever occurred first. The secondary endpoints are OS (time from registration to death from any cause), time to treatment failure (TTF: time from the enrollment date set as the start date until the day on which progression is confirmed, all-cause death occurs, or treatment is discontinued, whichever comes first), response rate (RR: proportion achieving complete or partial response), clinical benefit rate (CBR: complete response, partial response, or stable disease ≥ 6 months), and AEs. AEs are measured by the Common Terminology Criteria for Adverse Events (CTCAE) version 5.0.

Patients

Patients must meet all the following inclusion and exclusion criteria to be eligible as participants in this study.

Inclusion criteria were: 1) women ≥ 20 years old who are diagnosed with “histologically confirmed primary breast cancer”; 2) women whose breast cancer is HR+ HER2-; 3) women
who are diagnosed with “difficult to treat” MBC; 4) women
with a history of chemotherapy after diagnosis of MBC; 5)
women who plan to receive abemaciclib treatment; 6) women
who are scheduled to start abemaciclib treatment from 21 days
or later after completion of the previous chemotherapy and
14 days or later after completion of radiotherapy; 7) women
with no recent history of treatment with CDK4/6 inhibitors; 8)
women with no previous history of abemaciclib treatment; 9)
woman with an Eastern Cooperative Oncology Group (ECOG)
performance status (PS) of 0 - 2; 10) women whose vital or-
gan function is normal; and 11) women from whom written
informed consent to participate in the study is obtained.

Exclusion criteria were: 1) women for whom endocrine
therapy is not indicated; 2) women who require oxygen in-
halation because of dyspnea under resting conditions; 3)
women who have previous history of gastrectomy or small
bowel resection, except for mucosal resection; 4) women who
have Crohn’s disease, ulcerative colitis, or chronic diarrhea of
grade 2 or higher; 5) women who are pregnant or nursing; 6)
women undergoing treatment for bacterial or fungal infection;
7) women who have active hepatitis B or C; 8) women who
have serious cardiac disease; and 9) women whose primary
care physicians considered that they are unsuitable for inclu-
sion in this study.

Sample size

Since the PALOMA3 trial is the closest approximation to our
current study (in terms of containing patients that had received
prior chemotherapy), we have used PFS from that study as our
benchmark. Hence, the expected median PFS in the current
clinical trial is set at 7.7 months.

Meanwhile, in studies investigating efficacy of palboci-
clib for chemotherapy-treated patients, the median PFS was
reported to be 5.9 months in patients treated with one or two
regimens, and 4.3 months in patients treated with three regi-
mens or more [14]. As prognosis therefore appears to correlate
inversely with number of treatment regimens, we expect a me-
dian PFS of 7.7, 5.9, and 4 months in patients treated with one,
two, or three or more regimens, respectively.

In this study, it is important to verify that the median
PFS is 6 months or longer in patients previously treated with
a single regimen chemotherapy. Therefore, we will include a
sample size that can be used to determine the median PFS in
these patients with high accuracy. If expected median PFS is 8
months under conditions where 160 patients are enrolled with-
in 24 months of the enrollment period and within 36 months
of the total study period, the probability that 95% confidence
interval (CI) falls into less than 4 months is 80%; the prob-
ability rises to 90% if 95% CI falls into less than 4.5 months.
We will therefore set the sample size at 300 to ensure that at
least 160 patients will have been treated with a single regimen
of chemotherapy by the end of this study. Under these condi-
tions, we can accurately determine the median PSF in patients
treated with two or more regimens of chemotherapy and in
those receiving maintenance therapy.

Discontinuation of subjects

Under the following conditions, participation in the study
by subjects from whom consent has been obtained can be
discontinued. 1) When subjects withdraw their consent. 2)
When the principal investigator decides that withdrawal is appropriate.

**Treatment methods**

In combination with endocrine agents, abemaciclib (150 mg) is usually administered orally twice daily to an adult. Dosage should be reduced according to the condition of each patient as appropriate. There will be no provision on endocrine agents to be used in combination with abemaciclib. This will be determined through consultation between the attending physician and the patient as part of routine clinical practice.

**Data collection**

Clinical data are prospectively obtained from the records generated in each institution. In this study, the personnel will enter and correct data using Research Electronic Data Capture (REDCap).

**Data management**

In the REDCap system, revision records of data correction will be recorded in a log file. Other than for data monitoring, data access for quality management is granted to the director of data management in this study.

**Statistical analysis**

All patients who receive abemaciclib and who meet the inclusion criteria will be included in the analysis. The median values of PFS, OS, and TTF will be calculated. Survival curves will be calculated using the Kaplan-Meier method. Following this, the 1-, 2-, and 3-year survival rates and corresponding 95% CIs will be calculated. The number and proportion of patients showing response or clinical benefit in the efficacy analysis set will be calculated. The corresponding 95% CIs will also be calculated. Compliance will be calculated for both hormonal therapy and abemaciclib treatment. The highest grade of AEs observed for individual patients will be tallied.

Subpopulation analysis will be conducted to determine whether there is any difference in PFS among subpopulations. The preplanned subpopulation analysis is the number of chemotherapy regimens for HR+ HER2- MBC (two or less vs. three or more), prior treatment history with CDK4/6 inhibitors other than abemaciclib (presence vs. absence) and menopausal status (pre vs. post).

We also planned to determine PFS of the subpopulation treated with abemaciclib as maintenance therapy after chemotherapy. Maintenance therapy in this study is defined as follows: the former chemotherapy was performed more than 9 weeks, the best response of the former chemotherapy was determined as CR, PR or SD, and switching to abemaciclib therapy was planned for a reason other than PD.

**Ethics approval**

This study has been approved by the Ethical Review Board of Yokohama City University on November 28, 2019 (approval number: B191000019) and registered at the UMIN Clinical Trials Registry as UMIN 000037395. Ethical approval was obtained from all institutes involved in CSPOR-BC. The study will be conducted in accordance with the legal and regulatory requirements as well as the general principles set forth in the International Ethical Guidelines for Biomedical Research Involving Human Subjects (Council for International Organizations of Medical Sciences 2002), Guidelines of Good Clinical Practice (International Conference on Harmonization 1996), and the Declaration of Helsinki (World Medical Association 1996 and 2008).

**Discussion**

We started a prospective cohort study to verify the clinical efficacy and safety of combination therapy with abemaciclib and hormone therapy in patients with MBC who underwent chemotherapy based on the hypothesis that this combination therapy may be effective even after chemotherapy. Unlike other CDK4/6 inhibitors, abemaciclib causes neutropenia at a low frequency, thus it could be administered safely and is more acceptable in chemotherapy-treated patients. Additionally, abemaciclib has been reported to have a fairly good outcome in chemotherapy-treated patients with MBC in the MONARCH-1 trial with an objective response rate of 19.7%, a CBR of 42.4%, and a median PFS of 6.0 months [15].

Notably, there are other issues serving as reasons for evaluating abemaciclib combination therapy for chemotherapy-treated patients. One issue is that treatment with a CDK4/6 inhibitor followed by another CDK4/6 inhibitor is not recommended because of the limited data to support it [12]. Basic research has reported the cross resistance of CDK4/6 inhibitors. However, recovery of sensitivity after a temporary cessation of exposure has also been reported [16]. In this regard, it may be better to use a different mechanism after administration of an initial CDK4/6 inhibitor before using another CDK4/6 inhibitor. There is another report on recovery from drug resistance by improving microenvironmental hypoxia and epithelial-mesenchymal transition after the administration of eribulin [17]. Therefore, initial CDK4/6 inhibitor administration followed by chemotherapy then abemaciclib treatment could be one of the options for MBC treatment whose safety and efficacy are worth verifying.

On the other hand, there are some patients requiring more effective treatment from the diagnosis of HR+ HER2- MBC. These patients require chemotherapy from the start of their treatment because of their life-threatening disease or severe symptoms such as pain or discharge from the exposed lesion. Although a favorable response is obtained, it occasionally appears better to discontinue the initial chemotherapy owing to AEs such as cardiotoxicity caused by anthracyclines and peripheral neuropathy caused by taxanes. The combination of CDK4/6 inhibitors and hormone therapy has the potential to be a practical therapy as maintenance therapy after chemotherapy.
The clinical efficacy and safety of this combination under such condition will be verified.

In conclusion, this prospective study aims to investigate the efficacy and safety of combination therapy with endocrine therapy and abemaciclib in patients who have treatment history with chemotherapeutics for MBC in daily clinical practice.

**Trial status**

This study started recruiting patients from December 2019 and will continue recruitment until November 2022 from designated hospitals in Japan. The protocol version is 2.0, which was established on July 8, 2021.

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**Conflict of Interest**

The authors declare that they have no conflict of interest.

**Informed Consent**

All eligible patients granted written informed consent.

**Author Contributions**

All authors conceived and designed. KN and KM conceived, designed, interpreted data, and drafted the manuscript. KA assisted an ancillary study for drug-induced lung injury caused by abemaciclib and paper revisions. YU completed analyses for trial data. HM supervised the overall study and approved the submission of the study design to the journal as the representative of CSPOR-BC. All authors read and approved the manuscript.

**Data Availability**

Data supporting the trial will be made available on the CSPOR-BC website (https://cspor-bc.or.jp). Detailed datasets for this trial will be kept in the data center of the CSPOR-BC office and will not be accessible during data collection. At trial conclusion, summary data will be made available on the website to aid interpretation and replication of analyses.

**Author Note**

The authors belong to CSPOR-BC, a clinical trial group studying breast cancer in Japan since 2000. The authors have managed and disseminated the data of various clinical trials from Japan.

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