Phase II study of neoadjuvant anthracycline combined with nanoparticle albumin-bound paclitaxel for human epidermal growth factor receptor 2-negative breast cancer

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Key words: nanoparticle albumin-bound paclitaxel, neoadjuvant chemotherapy, human epidermal growth factor receptor 2-negative breast cancer, pathological complete response, clinical trial, phase II therapy. Thus, neoadjuvant therapy with FEC followed by nab-PTX for operable HER2-negative breast cancer was found to be a safe and effective option.

Introduction

Neoadjuvant chemotherapy (NAC) with anthracyclines followed by taxane chemotherapy has been widely recognized as a standard therapy for patients with locally advanced breast cancer. In recent years, this approach has been adopted for the treatment of not only locally advanced, but also resectable breast cancer (1-3). The rate of breast-conserving surgery (BCS) is increased by NAC (4). Regimens including 5-fluorouracil, epirubicin and cyclophosphamide (FEC), epirubicin and cyclophosphamide (EC), or doxorubicin and cyclophosphamide (AC) have been selected as the anthracycline regimens (2,3,5-7), whereas docetaxel or paclitaxel (PTX) have been used as the taxane regimens (8-13). Nanoparticle albumin-bound PTX (nab-PTX) has recently attracted attention as a treatment for advanced metastatic or recurrent breast cancer (14,15).

Nab-PTX is an amorphous protein product made from solvent-free PTX with human albumin. Nab-PTX was developed to decrease toxicity and increase antitumor activity, and it does not require steroid administration to prevent hypersensitivity reactions.

A phase III trial on metastatic breast cancer comparing standard PTX 175 mg/m^2 every 3 weeks with nab-PTX 260 mg/m^2 every 3 weeks demonstrated significant improvement in the response rate (19 vs. 33%, respectively; P=0.001) and in the interval to progression (16.9 vs. 23.0 weeks, respectively) in the nab-PTX arm, with a hazard ratio of 0.75 (P=0.006) (14).

Data on neoadjuvant regimens including nab-PTX for HER2-negative breast cancer are limited. To the best of our knowledge, a trial of FEC followed by nab-PTX has not yet been reported. Thus, a phase II trial was undertaken to evaluate the safety and effectiveness of this regimen.

Patients and methods

Patients. This was a single-center, prospective, single-arm, phase II trial. Patients with histologically confirmed invasive
breast cancer by core biopsy with lymph node-positive, HER2-negative, operable invasive breast cancer (clinical stage T1-4N1-3) were considered eligible. All the tumors were evaluated locally for expression of estrogen receptor (ER), progesterone receptor (PgR) and HER2. ER and PgR were defined as positive with positive staining of >1%. HER2 was defined as positive with either positivity on immunohistochemistry of 3+, or demonstration of amplification on fluorescence in situ hybridization of >2. Other eligibility criteria included Eastern Cooperative Oncology group (EGOG) performance status ≤1; age ≥20 years; hemoglobin level ≥8.0 g/dl; white blood cell count 3,000-12,000/µm³; platelet count ≥10x10⁹/µm³; creatinine level ≤1.5 mg/dl; normal total liver function, with bilirubin ≤1.5 mg/dl, aspartate aminotransferase <100 IU/l and alanine aminotransferase <100 IU/l; and normal cardiac function.

Patients who had received any previous treatment were excluded. Other exclusion criteria included severe heart disease, diarrhea, ileus, intestinal paralysis, poorly controlled diabetes, poorly controlled angina, myocardial infarction, heart failure within 6 months, interstitial lung fibrosis, cerebrovascular disorder, symptomatic brain metastases, gastrointestinal bleeding requiring blood transfusion, severe bone marrow suppression, severe renal disorder, severe liver disorder, severe pleural effusion, severe ascites, previous hypersensitivity to nab-PTX or albumin, pentostatin treatment, other active cancers, any infectious disease, pregnancy or lactation, mental disorder, or steroid therapy.

This trial was approved by the Ethics Committee of the Chiba University Graduate School of Medicine and all the patients provided written informed consent prior to inclusion (registered at UMIN000007724).

**Therapy.** The patients received 4 cycles of FEC (5-fluorouracil 500 mg/m², epirubicin 100 mg/m² and cyclophosphamide 500 mg/m² every 21 days) followed by 4 cycles of nab-PTX at 260 mg/m² every 21 days. Adverse events were graded according to the National Cancer Institute Common Terminology Criteria for Adverse Events, version 4.0 (https://evs.nci.nih.gov/ftp1/CTCAE/CTCAE_4.03_2010-06-14_QuickReference_5x7.pdf). Preventive granulocyte colony-stimulating factor was not administered. The patients underwent surgery 4-6 weeks after the chemotherapy.

**Endpoint.** The primary endpoint was pathological complete response (pCR) rate, defined as no histological residual invasive tumor cells in the breast and axillary lymph nodes (ypT0/Tis and ypN0). There was no case of ypT0/Tis and ypN0.

The secondary endpoints were clinical response rate, BCS rate and safety. Clinical response rates were evaluated by ultrasonography, computed tomography and magnetic resonance imaging according to the Response Evaluation Criteria in Solid Tumors guidelines, version 1.1 (https://ctep.cancer.gov/protocoldevelopment/docs/recist_guideline.pdf).

**Statistical analysis.** Robidoux et al reported that the phase II trial National Surgical Adjuvant Breast and Bowel Project (NSABP) FB-AX-003 of neoadjuvant weekly nab-PTX followed by FEC for locally advanced breast cancer demonstrated a pCR rate of 17% in the HER2-negative subset (16). In another previous study, NSABP B-27 of AC followed by docetaxel demonstrated a pCR rate of 26% (3). For the present study, a threshold pCR rate of 10%, an expected pCR rate of 25%, an α error of 5%, and a power of 80% were set based on these previous studies. Consequently, an accrual of 36 patients was planned to produce a minimum of 33 evaluable patients.

**Results**

**Patient population.** Between November 2011 and October 2013, 19 patients were enrolled; 3 patients were excluded due to withdrawal of consent prior to treatment initiation and 16 patients were finally considered evaluable for efficacy and safety of this study treatment.

The patients' characteristics are summarized in Table I. The median age was 54 years, 3 patients had stage II breast cancer, and 13 patients had stage III breast cancer. Of the 16 patients, 8 (50%) had luminal type B breast cancer, and 8 (50%) had triple-negative breast cancer.

**Treatment administration and study completion.** A total of 10 (62%) patients received all 8 planned cycles without delays or missed doses; 3 (18%) patients missed doses and/or required delays during FEC therapy due to adverse events; 4 patients missed doses (185-220 mg/m²) during the first administration of nab-PTX.

**Clinical and pathological assessments.** The pCR rate was 18.75% (3/16 patients); 12.5% (1/8 patients) had luminal type B and 25% (2/8 patients) had triple-negative breast cancer. The clinical response rate was 100% (clinical complete and partial response in 6 and 10 patients, respectively) (Table II). The pathological treatment effect was as follows: Grade 1a, n=8; grade 1b, n=3; grade 2a, n=1; grade 2b, n=1; and grade 3, n=3 (Table III). The BCS rate was 31.25%.

**Safety profile.** The adverse events are listed in Table IV. A total of 3 patients developed grade 3 neutropenia. Grade 3 adverse events occurred during FEC therapy, but were not observed during nab-PTX therapy. Nausea and vomiting occurred more frequently during FEC.

All the patients experienced grade 1/2 muscle pain and peripheral neuropathy during nab-PTX treatment. Grade 3/4 non-hematological toxicity did not occur with either regimen. No unexpected severe adverse events occurred.

**Discussion**

To the best of our knowledge, this is the first report of a phase II study of neoadjuvant nab-PTX with the FEC regimen for HER2-negative breast cancer.

High pCR rates with neoadjuvant regimens have been reported for HER2-positive breast cancer (17), but not for HER2-negative breast cancer. In the NSABP FB-AX-003 phase II trial, the pCR rate for HER2-positive cancer with a 24-week regimen of weekly nab-PTX plus trastuzumab, followed by FEC and trastuzumab, was 58%, whereas the pCR rate of the regimen without trastuzumab for HER2-negative cancer was 17% (16). Nab-PTX has been shown to be effective in...
Neoadjuvant FEC followed by nab-PTX has not yet been reported. Thus, the safety and effectiveness of this regimen as neoadjuvant chemotherapy for operable HER2-negative breast cancer needed to be assessed. Thus, a single-center, phase II study was performed to evaluate the safety and pCR of this regimen. In our hospital, the pCR rate for AC followed by tri-weekly PTX was 5.9%, for FEC followed by tri-weekly PTX it was 13.2%, and for FEC followed by nab-PTX it was 18.75%. Shimada et al. reported that the pCR rate for nab-PTX followed by EC was 5.7% (18). Two explanations were considered as plausible for the low pCR rate in that study: First, FEC therapy may be more effective compared with EC therapy. Second, 64% of patients in that study had ER-positive breast cancer, which may have included some patients with luminal type A breast cancer. In the present study, 50% of the patients had ER-positive breast cancer.

Of note, in the present study, the clinical response rate was 100%. Shimada et al. reported that nab-PTX followed by EC achieved a clinical response rate of 75.5% (18). We considered that the reason for the lower clinical response rate in that study was also the difference between FEC and EC therapy. Second, that study included several patients with ER-positive breast cancer (14,15). Neoadjuvant FEC followed by nab-PTX has not yet been reported.
breast cancer. The efficacy of FEC followed by nab-PTX as neoadjuvant chemotherapy for patients with HER2-negative operable cancer may be higher compared with that of methods used to date when the pCR rate is evaluated.

Three patients (18.75%) had grade 3–4 AEs, and all patients had grade 1–2 AEs. During nab-PTX treatment, no patient had grade 3–4 AEs. Thus, FEC did not appear to adversely affect nab-PTX therapy. A previous trial of AC followed by PTX reported that grade 3–4 AEs during PTX included neurosensory toxicity in 15%, neuromotor toxicity in 7%, neurosensory or neuromotor toxicity in 18%, arthralgia and/or myalgia in 12%, day 1 granulocytopenia in 3%, and febrile neutropenia in 3% of the patients (12). Another previous trial of AC followed by PTX reported grade 4 neutropenia in 16% during PTX therapy, and 3% patients required hospitalization due to AEs during PTX therapy (11). They reported that a severe hypersensitivity reaction occurred in 1% of patients during PTX administration (11). However, there were no grade ≥3 AEs during nab-PTX in the present trial. All the patients had grade 1-2 AEs, but the FEC regimen followed by nab-PTX appeared to have good tolerability. The present study was prematurely terminated due to its poor accrual rate. The time to enroll patients was not extended, as we had started another neoadjuvant trial including nab-PTX for patients with HER2-negative cancer.

In conclusion, FEC followed by nab-PTX as neoadjuvant treatment for HER2-negative operable breast cancer appeared to be effective and well-tolerated. This regimen exhibited better efficacy compared with the standard anthracycline chemotherapy regimen, whereas, in terms of side effects, this combination may be among the safest.

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References