

- radiation-induced DNA damage in lymphocytes. *Breast Cancer Res* 2005;7:R690–R698.
19. Chang-Claude J, Popanda O, Tan XL, *et al.* Association between polymorphisms in the DNA repair genes, XRCC1, APE1, and XPD and acute side effects of radiotherapy in breast cancer patients. *Clin Cancer Res* 2005;11:4802–4809.
 20. Andreassen CN, Alsner J, Overgaard M, *et al.* Prediction of normal tissue radiosensitivity from polymorphisms in candidate genes. *Radiother Oncol* 2003;69:127–135.
 21. Angele S, Romestaing P, Moullan N, *et al.* ATM haplotypes and cellular response to DNA damage: association with breast cancer risk and clinical radiosensitivity. *Cancer Res* 2003;63:8717–8725.
 22. Quarmby S, Fakhoury H, Levine E, *et al.* Association of transforming growth factor beta-1 single nucleotide polymorphisms with radiation-induced damage to normal tissues in breast cancer patients. *Int J Radiat Biol* 2003;79:137–143.
 23. Andreassen CN, Alsner J, Overgaard J, *et al.* TGFB1 polymorphisms are associated with risk of late normal tissue complications in the breast after radiotherapy for early breast cancer. *Radiother Oncol* 2005;75:18–21.
 24. Andreassen CN, Overgaard J, Alsner J, *et al.* ATM sequence variants and risk of radiation-induced subcutaneous fibrosis after postmastectomy radiotherapy. *Int J Radiat Oncol Biol Phys* 2006;64:776–783.
 25. Giotopoulos G, Symonds RP, Foweraker K, *et al.* The late radiotherapy normal tissue injury phenotypes of telangiectasia, fibrosis and atrophy in breast cancer patients have distinct genotype-dependent causes. *Br J Cancer* 2007;96:1001–1007.
 26. Andreassen CN, Alsner J, Overgaard M, *et al.* Risk of radiation-induced subcutaneous fibrosis in relation to single nucleotide polymorphisms in TGFB1, SOD2, XRCC1, XRCC3, APEX and ATM—A study based on DNA from formalin fixed paraffin embedded tissue samples. *Int J Radiat Biol* 2006;82:577–586.
 27. Iwakawa M, Noda S, Ohta T, *et al.* Different radiation susceptibility among five strains of mice detected by a skin reaction. *J Radiat Res* 2003;44:7–13.
 28. Ohta T, Iwakawa M, Oohira C, *et al.* Fractionated irradiation augments inter-strain variation of skin reactions among three strains of mice. *J Radiat Res* 2004;45:515–519.
 29. Iwakawa M, Noda S, Ohta T, *et al.* Strain dependent differences in a histological study of CD44 and collagen fibers with expression analysis of inflammatory response-related genes in irradiated murine lung. *J Radiat Res* 2004;45:423–433.
 30. Noda S, Iwakawa M, Ohta T, *et al.* Inter-strain variance in late phase of erythematous reaction or leg contracture after local irradiation among three strains of mice. *Cancer Detect Prev* 2005;29:376–382.
 31. Ishikawa K, Koyama-Saegusa K, Otsuka Y, *et al.* Gene expression profile changes correlating with radioresistance in human cell lines. *Int J Radiat Oncol Biol Phys* 2006;65:234–245.
 32. Ban S, Ishikawa K, Kawai S, *et al.* Potential in a single cancer cell to produce heterogeneous morphology, radiosensitivity and gene expression. *J Radiat Res* 2005;46:43–50.
 33. Tsuji AB, Sudo H, Sugyo A, *et al.* A fast, simple method for screening radiation susceptibility genes by RNA interference. *Biochem Biophys Res Commun* 2005;333:1370–1377.
 34. Bennett CB, Lewis LK, Karthikeyan G, *et al.* Genes required for ionizing radiation resistance in yeast. *Nat Genet* 2001;29:426–434.
 35. Rouse J, Jackson SP. Interfaces between the detection, signaling, and repair of DNA damage. *Science* 2002;297:547–551.
 36. Kolodner RD, Putnam CD, Myung K. Maintenance of genome stability in *Saccharomyces cerevisiae*. *Science* 2002;297:552–557.
 37. Nasmyth K. Segregating sister genomes: the molecular biology of chromosome separation. *Science* 2002;297:559–565.
 38. Iwakawa M, Noda S, Yamada S, *et al.* Analysis of non-genetic risk factors for adverse skin reactions to radiotherapy among 284 breast cancer patients. *Breast Cancer* 2006;13:300–307.
 39. Abecasis GR, Cookson WO. GOLD—Graphical overview of linkage disequilibrium. *Bioinformatics* 2000;16:182–183.
 40. Zhao JH, Lissarrague S, Essioux L, *et al.* GENECOUNTING: Haplotype analysis with missing genotypes. *Bioinformatics* 2002;18:1694–1695.
 41. Clark AG. The role of haplotypes in candidate gene studies. *Genet Epidemiol* 2004;27:321–333.
 42. Fallin D, Cohen A, Essioux L, *et al.* Genetic analysis of case/control data using estimated haplotype frequencies: Application to APOE locus variation and Alzheimer's disease. *Genome Res* 2001;11:143–151.
 43. Hirakawa M, Tanaka T, Hashimoto Y, *et al.* JSNP: a database of common gene variations in the Japanese population. *Nucleic Acids Res* 2002;30:158–162.
 44. Smigielski EM, Sirotkin K, Ward M, *et al.* dbSNP: A database of single nucleotide polymorphisms. *Nucleic Acids Res* 2000;28:352–355.
 45. Buetow KH, Edmonson M, MacDonald R, *et al.* High-throughput development and characterization of a genomewide collection of gene-based single nucleotide polymorphism markers by chip-based matrix-assisted laser desorption/ionization time-of-flight mass spectrometry. *Proc Natl Acad Sci U S A* 2001;98:581–584.
 46. Kamatani N, Kawamoto M, Kitamura Y, *et al.* Establishment of B-cell lines derived from 996 Japanese individuals. *Tissue Cult Res Commun* 2004;23:71–80.
 47. Singer VL, Jones LJ, Yue ST, Haugland RP. Characterization of PicoGreen reagent and development of a fluorescence-based solution assay for double-stranded DNA quantitation. *Anal Biochem* 1997;249:228–238.
 48. Schaid DJ, Rowland CM, Tines DE, *et al.* Score tests for association between traits and haplotypes when linkage phase is ambiguous. *Am J Hum Genet* 2002;70:425–434.
 49. Barrett JC, Fry B, Maller J, *et al.* Haploview: Analysis and visualization of LD and haplotype maps. *Bioinformatics* 2005;21:263–265.
 50. Storey JD, Tibshirani R. Statistical significance for genomewide studies. *Proc Natl Acad Sci U S A* 2003;100:9440–9445.
 51. Benjamini Y, Yekutieli D. The control of the false discovery rate in multiple testing under dependency. *Ann Stat* 2001;29:1165–1188.
 52. Zhang K, Zhu J, Shendure J, *et al.* Long-range polony haplotyping of individual human chromosome molecules. *Nat Genet* 2006;38:382–387.
 53. Aruffo A, Stamenkovic I, Melnick M, *et al.* CD44 is the principal cell surface receptor for hyaluronate. *Cell* 1990;61:1303–1313.
 54. Weinfeld M, Rasouli-Nia A, Chaudhry MA, *et al.* Response of base excision repair enzymes to complex DNA lesions. *Radiat Res* 2001;156:584–589.
 55. Dianov GL, O'Neill P, Goodhead DT. Securing genome stability by orchestrating DNA repair: Removal of radiation-induced clustered lesions in DNA. *Bioessays* 2001;23:745–749.
 56. Hang H, Rauth SJ, Hopkins KM, *et al.* Mutant alleles of *Schizosaccharomyces pombe* Rad9(+) alter hydroxyurea resistance, radioresistance and checkpoint control. *Nucleic Acids Res* 2000;28:4340–4349.
 57. Komatsu K, Miyashita T, Hang H, *et al.* Human homologue of *S. pombe* Rad9 interacts with BCL-2/BCL-x_L and promotes apoptosis. *Nat Cell Biol* 2000;2:1–6.
 58. International HapMap Consortium. A haplotype map of the human genome. *Nature* 2005;437:1299–1320.
 59. Ashburner M, Ball CA, Blake JA, *et al.*, for the Gene Ontology Consortium. Gene ontology: Tool for the unification of biology. *Nat Genet* 2000;25:25–29.

Dofequidar Fumarate (MS-209) in Combination With Cyclophosphamide, Doxorubicin, and Fluorouracil for Patients With Advanced or Recurrent Breast Cancer

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A B S T R A C T

Purpose

To evaluate the efficacy and tolerability of dofequidar plus cyclophosphamide, doxorubicin, and fluorouracil (CAF) therapy in comparison with CAF alone, in patients with advanced or recurrent breast cancer. Dofequidar is a novel, orally active quinoline derivative that reverses multidrug resistance.

Patients and Methods

In this randomized, double-blind, placebo-controlled trial, patients were treated with six cycles of CAF therapy: 28 days/cycle, with doxorubicin (25 mg/m²) and fluorouracil (500 mg/m²) administered on days 1 and 8 and cyclophosphamide (100 mg orally [PO]) administered on day 1 through 14. Patients received dofequidar (900 mg PO) 30 minutes before each dose of doxorubicin. Primary end point was overall response rate (ORR; partial or complete response). In total, 221 patients were assessable.

Results

ORR was 42.6% for CAF compared with 53.1% for dofequidar + CAF, a 24.6% relative improvement and 10.5% absolute increase ($P = .077$). There was a trend for prolonged progression-free survival (PFS; median 241 days for CAF v 366 days for dofequidar + CAF; $P = .145$). In retrospectively defined subgroups, significant improvement in PFS in favor of dofequidar was observed in patients who were premenopausal, had no prior therapy, and were stage IV at diagnosis with an intact primary tumor. Except for neutropenia and leukopenia, there was no statistically significant excess of grade 3/4 adverse events compared with CAF. Treatment with dofequidar did not affect the plasma concentration of doxorubicin.

Conclusion

Dofequidar + CAF was well tolerated and is suggested to have efficacy in patients who had not received prior therapy.

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INTRODUCTION

Despite the advances in chemotherapeutic intervention, many cancers are either inherently resistant or develop resistance to chemotherapy.^{1,2} Consequently, multidrug resistance (MDR) remains a major obstacle to the successful treatment of cancer.^{1,3,4} One mechanism by which MDR operates is via the increased cellular efflux of cytotoxic compounds due to increased expression of membrane transport proteins such as P-glycoprotein (P-gp) and MDR-associated protein (MRP).^{1,4,5} MDR affects many structurally and functionally unrelated agents including cytotoxic drugs that are hydrophobic, natural products, such as taxanes, vinca alkaloids,

anthracyclines, epipodophyllotoxins, topotecan, dactinomycin, and mitomycin.^{1,6,7} These represent some of the most commonly used chemotherapeutic agents.

In tumors with low levels of P-gp expression at baseline or diagnosis, P-gp expression increases after exposure to chemotherapy agents, thus leading to the development of MDR. In breast cancer patients who had received prior chemotherapy, P-gp expression has been shown to increase from 11% in untreated patients to 30% after chemotherapy.⁸ Furthermore, compared with P-gp-negative tumors, a significant increase in resistance to paclitaxel and doxorubicin was reported in P-gp positive breast cancer tissue, irrespective of prior therapy.

The degree of P-gp expression also strongly correlated with the degree of drug resistance observed.⁸

Chemotherapy remains the treatment of choice for women with hormone receptor–negative and hormone-refractory breast cancer disease.^{9–11} However, many tumors that are initially responsive to chemotherapy frequently relapse and develop resistance to the broad spectrum of cytotoxic drugs currently employed.^{8,12,13} Consequently, MDR remains a major reason for treatment failure in patients with metastatic breast cancer and highlights the urgent need for MDR modifiers in breast cancer chemotherapy.

Since the discovery of verapamil as an MDR-reversing agent,¹⁴ many compounds have been investigated as MDR inhibitors.^{14–16} Dofequidar fumarate (Fig 1), is a novel, orally active, quinoline-derived inhibitor of MDR.¹⁷ In preclinical studies, dofequidar reversed MDR in P-gp– and MRP-1–expressing cancer cells in vitro (1 to 3 $\mu\text{mol/L}$), as well as enhancing the antitumor effects of doxorubicin in MDR tumor–bearing mice.^{17–19} A phase I trial in healthy volunteers showed dofequidar to be well tolerated (10 to 1,200 mg) with no dose-limiting toxicities and an effective plasma concentration was maintained for 8 hours at 900 mg (data on file, Schering AG, Berlin, Germany). In a phase II combination trial in patients with recurrent breast cancer, dofequidar potentiated the antitumor effects of CAF (cyclophosphamide, doxorubicin, and fluorouracil) therapy; patients who had not responded to treatment with three cycles of CAF responded to subsequent treatment with dofequidar plus CAF. The numbers of patients with an objective response were two of seven at 600 mg and two of six at 900 mg dofequidar, though dose escalation was stopped at 1,200 mg due to increased hematologic toxicity (data on file, Schering AG). On the basis of this result, this phase III study was conducted to compare the efficacy and safety of dofequidar plus CAF with placebo plus CAF in patients with advanced or recurrent breast cancer.

PATIENTS AND METHODS

Study Design

This was a randomized, multicenter, double-blind, placebo-controlled trial conducted at 46 centers across Japan, comparing the efficacy and safety of dofequidar plus CAF with placebo plus CAF. Female patients (age 20 to 70 years) with advanced (stage IV at diagnosis with an intact primary tumor) or recurrent breast cancer were enrolled onto the study. Other inclusion criteria included a histologically defined, measurable or assessable primary lesion; two or fewer regimens of prior chemotherapy in both neo/adjuvant and metastatic

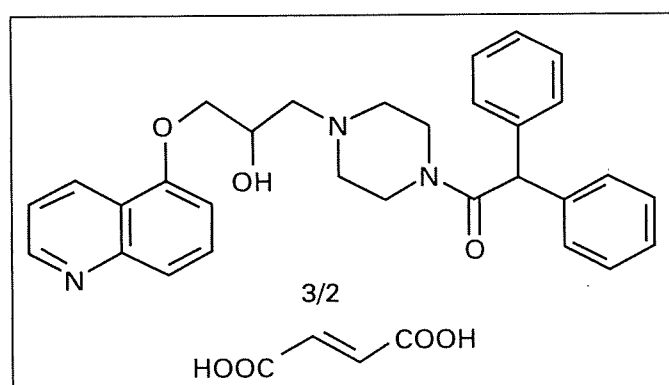


Fig 1. Structure of dofequidar (MS-209).

settings, (excluding prior endocrine or single-agent fluorouracil therapy); 180 mg/m^2 anthracyclines (doxorubicin equivalent) or less previously; a performance status of 0 to 2; and adequate bone marrow, renal, hepatic and cardiac functions. Patients who progressed or had a recurrence in less than 6 months with anthracycline-containing chemotherapy, and those who had a history of major cardiac disease, uncontrolled hypertension, symptomatic brain metastasis, or simultaneous malignancy were excluded. The trial was approved by the institutional review board and was conducted in accordance with the Declaration of Helsinki (1996). All patients provided written informed consent before study entry.

Dosing and Dose Modification for Toxicity

Patients were treated with six cycles of CAF therapy with dofequidar or placebo, and each treatment cycle lasted for 28 days; drugs were administered as follows: days 1 and 8, doxorubicin (25 mg/m^2) and fluorouracil (500 mg/m^2), each infused over 15 minutes; days 1 through 14, cyclophosphamide (100 mg orally [PO]); dofequidar (900 mg/d; 3 \times 300 mg tablets) or placebo administered 30 minutes before each doxorubicin dose to ensure adequate blood concentration of dofequidar. The doses of doxorubicin and fluorouracil were reduced to 20 mg/m^2 and 400 mg/m^2 , respectively, if any of the following criteria were met: grade 3 nonhematologic toxicity (except nausea and vomiting); grade 3 or worse neutropenia ($< 1,000/\text{mm}^3$) maintained for at least 5 days with an episode of fever of 38.5°C or higher; grade 3 or worse thrombocytopenia ($< 50,000/\text{mm}^3$); and grade 4 neutropenia ($< 500/\text{mm}^3$). The next cycle was postponed for 3 weeks unless the patient had a WBC count of at least 4,000/ mm^3 , or a neutrophil count of at least 2,000/ mm^3 and a platelet count of at least 100,000/ mm^3 . Patients were followed up for 3 months after completion or discontinuation of treatment.

Treatment Assignment

Patients were randomly assigned to their treatment by the Trial Register Center. Treatment assignment was securely stored and coded until completion of the study. Investigators were also blinded to the assigned treatment. Patients were stratified by the number of prior chemotherapy regimens, including adjuvant chemotherapy, by a history of prior use of anthracyclines, and by the presence of liver metastases.

Efficacy

The primary study end point was the overall response rate (ORR) in the full analysis set (FAS; all patients who received treatment at least once and met all inclusion/exclusion criteria). Efficacy assessment by lesion and ORR assessment were made at each treatment cycle (every 4 weeks) and at treatment completion. Objective responses were assessed through blinded reading of radiographs by an independent expert panel. The secondary study end points included complete response rate (CR), time to treatment failure (TTF), time to progression (TTP), and progression-free survival (PFS).

Subgroup analyses were conducted to assess PFS within specific patient subpopulations, including premenopausal women, patients who had no prior therapy, and patients who had advanced primary breast cancer.

Safety and Tolerability

Adverse events (AEs) were recorded at the end of each treatment cycle and at the end of the study period using data from the safety population (all patients who received treatment at least once in the study). AEs were categorized according to the National Cancer Institute Common Toxicity Criteria (NCI-CTC) Version 2. The incidence of significant decreases in left ventricular ejection fraction (LVEF) and serious AEs were recorded. The CBC was evaluated weekly. Serum chemistries and urinalysis were evaluated every 2 weeks. The minimum hematology values and LVEF in each treatment cycle were also recorded and analyzed in the per-protocol set (PPS; all patients who received treatment at least once and had no protocol deviations).

Pharmacokinetics

To assess the effect of concomitant dofequidar use on the pharmacokinetics of doxorubicin, the plasma doxorubicin concentration on day 1 of cycle 1 was compared between treatment groups. Blood samples were taken at baseline and at 15 minutes, 30 minutes, and 1, 2, 4, and 6 hours after the start of doxorubicin administration. Plasma doxorubicin concentrations were determined by reversed-phase high-performance liquid chromatography. Area

under the plasma concentration-time curve (AUC) was calculated using the linear trapezoidal rule.

Statistical Analyses

The primary end point was analyzed using the Fisher's exact test at a significance level of 2.5% in a one-sided test. A difference in response rates of 20% between the two treatment groups was used as the basis for a statistically significant difference. CR, TTF, TTP and PFS were analyzed by the log-rank test at a significance level of 5% in a two-sided test. The CR, TTF, TTP and PFS were analyzed in the FAS, and the pharmacokinetic data analyzed in the PPS.

RESULTS

Patient Characteristics

A total of 227 patients were recruited onto the study (Fig A1, online only), of which 225 patients were included in the safety analysis ($n = 113$ for the dofequidar group; $n = 112$ for the placebo group); two patients did not receive the study treatment and were thus excluded. Four patients did not meet the inclusion/exclusion criteria; therefore, the FAS consisted of 221 patients ($n = 113$ for the dofequidar group; $n = 108$ for the placebo group). The PPS consisted of 199 patients ($n = 100$ for the dofequidar group; $n = 99$ for the placebo group). There were 22 patients excluded from the PPS analysis due to protocol deviations. Baseline patient characteristics were well balanced between the two treatment arms (Table 1). Most patients had predominantly recurrent disease and had received prior chemotherapy plus endocrine therapy. Also, many patients who had advanced primary breast cancer had received no prior therapy.

Table 1. Patient Demographics (full analysis set)

Characteristic	Dofequidar + CAF ($n = 113$)		Placebo + CAF ($n = 108$)	
	No.	%	No.	%
Age, years				
Mean	54.4		52.4	
SD	7.69		8.97	
Medical history known	65	57.5	60	55.6
Weight, kg				
Mean	56.2		54.1	
SD	7.52		7.73	
Height, cm				
Mean	154.7		154.7	
SD	5.71		5.61	
Body surface area, m ²				
Mean	1.5		1.5	
SD	0.11		0.11	
Disease state				
Recurrent	81	71.7	80	74.1
Advanced	32	28.3	28	25.9
Prior therapy				
Radiotherapy + chemotherapy + endocrine therapy	32	22.1	32	29.6
Chemotherapy + endocrine therapy	55	48.7	54	50.0
Radiotherapy	1	0.9	1	0.9
No prior therapy	25	22.1	21	19.4
Menopausal status				
Premenopausal	24	21.2	26	24.1
Postmenopausal	88	77.9	79	73.1

Abbreviations: CAF, cyclophosphamide, doxorubicin, and fluorouracil; SD, standard deviation.

Efficacy

The ORR, rated as CR or partial response rate, was 42.6% for CAF plus placebo versus 53.1% for dofequidar plus CAF (Table 2). Although this represents a 24.6% relative improvement and a 10.5% absolute increase in response rate for patients receiving dofequidar plus CAF compared with those receiving CAF plus placebo, this response was not statistically significant ($P = .077$). A higher value was observed in the dofequidar treatment group for all secondary end points compared with placebo, though these results were not statistically significant. Among them, Figure 2 shows a trend for prolonged PFS (median, 241 days for CAF plus placebo v 366 days for dofequidar plus CAF; $P = .145$).

Dofequidar plus CAF significantly improved PFS in several patient subgroups, including patients who were premenopausal ($P = .046$; Fig 3A), patients who had not received prior therapy ($P = .0007$; Fig 3B), and patients who had advanced primary breast cancer ($P = .017$; Fig 3C). An extended follow-up showed that dofequidar plus CAF also significantly improved overall survival ($P = .0034$; Fig 3D) in patients who had no prior therapy.

Safety and Tolerability

A similar number of patients completed six treatment cycles in both groups ($n = 53$ for the dofequidar group; $n = 51$ for the placebo group). The mean number of treatment cycles was 4.5 in the dofequidar group and 4.3 in the placebo group. More than half of patients in both groups included in each cycle from cycle 2 onward had a delay in treatment, mostly due to prolonged hematologic toxicities.

Dofequidar plus CAF was well tolerated throughout the study. No statistically significant excess of grade 3/4 AEs, except for neutropenia ($P = .006$) and leukopenia ($P = .005$), was found in the dofequidar group compared with placebo (Table A1, online only). Importantly, there was no marked difference in the incidence of neutropenia-related morbidity, such as febrile neutropenia or infection, between the two treatment groups. No significant differences in the incidence of cardiac AEs were found between the two treatment groups. In addition, dose intensities of chemotherapeutic agents were similar in both treatment arms. No significant difference in the incidence of serious AEs (SAEs) was observed between either group. However, there was a trend for a higher incidence of SAEs from leukopenia in the dofequidar group than in the placebo group ($P = .060$; Fisher's exact test); five leukopenia cases were reported for dofequidar, whereas no such case was reported for placebo.

A total of 124 patients discontinued the study ($n = 61$ for the dofequidar group; $n = 63$ for the placebo group). The major reasons for discontinuation were progressive disease ($n = 23$ for the dofequidar group; $n = 28$ for the placebo group), grade 4 hematologic toxicity ($n = 20$ for the dofequidar group; $n = 6$ for the placebo group), failure to meet treatment continuation criteria ($n = 6$ for the dofequidar group; $n = 8$ for the placebo group), and consent withdrawal ($n = 6$ for the dofequidar group; $n = 12$ for the placebo group). Of the 225 patients who received treatment in the study, 14 patients died during the treatment period ($n = 3$), the follow-up period ($n = 2$), or the follow-up period after study termination ($n = 9$). There were 49 other serious AEs in 32 patients during the study and follow-up period.

Pharmacokinetics

The mean plasma concentrations of doxorubicin in the dofequidar- and placebo-treatment groups at 15 minutes postadministration reached 0.997 $\mu\text{g/mL}$ and 1.259 $\mu\text{g/mL}$, respectively, followed by biphasic elimination in both treatment groups. Mean plasma concentrations in

Table 2. Response Rates for Patients Treated With Dofequidar Plus CAF (n = 113) or Placebo Plus CAF (n = 108)

Treatment Group	Parameter (No. of patients)					Overall Response Rate (%)	95% CI
	Complete Response	Partial Response	No Change (stable disease)	Progressive Disease	Not Assessable		
Dofequidar	5	55	40	10	3	53.1	43.5 to 62.5
Placebo	4	42	41	14	7	42.6	33.1 to 52.5

NOTE. Odds ratio = 1.53 (range, 0.87-2.69); $P = .077$ for dofequidar v placebo. Abbreviation: CAF, cyclophosphamide, doxorubicin, and fluorouracil.

the dofequidar and placebo groups remained similar at 1, 2, 4, and 6 hours after the start of doxorubicin administration. Thus the elimination pattern for the first 6 hours after the start of administration was similar in both groups. The plasma concentrations of doxorubicin in the terminal phase (4 and 6 hours postadministration) were slightly higher in the dofequidar group compared with placebo (1.2- to 1.3-fold). However, AUC (0 to 6 hours) values showed no statistically significant difference between the dofequidar and placebo groups (mean, $0.480 \mu\text{g} \cdot \text{h/mL}$; standard deviation [SD], 0.324; range, 0.237-1.692; and mean, $0.407 \mu\text{g} \cdot \text{h/mL}$; SD, 0.062; and range, 0.289-0.500, respectively). Therefore, treatment with dofequidar did not affect the plasma concentrations of doxorubicin in patients (Fig 4).

DISCUSSION

Chemotherapy remains the preferred adjuvant treatment for patients with hormone receptor–negative disease and for patients with more aggressive, hormone receptor–positive tumors.^{11,20} However, despite the use of conventional adjuvant chemotherapy regimens, a significant proportion of patients with breast cancer still experience disease recurrence because of inherent or acquired drug resistance.¹² In this randomized phase III trial, the efficacy and safety of the multidrug resistance inhibitor dofequidar plus CAF was compared with CAF plus placebo in patients with recurrent or advanced breast cancer. Although, there was an observed relative improvement and absolute

increase in response rate for patients who received dofequidar plus CAF, these results did not reach statistical significance. This improvement in response rate may have been reflected in the observation that there was a trend for prolonged PFS, which favored patients in the dofequidar plus CAF group.

To date, only two randomized trials have examined the efficacy of a P-gp inhibitor in combination with chemotherapy in breast cancer patients. Wishart et al²¹ examined quinidine combined with epirubicin in patients with advanced breast cancer, but failed to show any significant difference in overall survival or PFS compared with placebo. In a more recent prospective study of patients with anthracycline-resistant metastatic breast cancer (n = 99), verapamil combined with vindesine and fluorouracil resulted in a significantly longer overall survival and a higher response rate compared with patients who did not receive the P-gp inhibitor (median survival, 323 v 209 days; $P = .036$, respectively; ORR, 27% v 11%; $P = .04$, respectively).²²

In the subgroup analyses, dofequidar in combination with CAF displayed a significantly increased PFS in patients who had not received prior therapy, who had advanced primary breast cancer or who were premenopausal. In addition, dofequidar also significantly improved overall survival in the patient group who had no prior therapy. Although the patient numbers in these analyses were small, the results remain important within these clinically significant patient populations. Both preclinical and clinical data have indicated that newer-generation MDR modulators can prevent the development of resistance.^{23,24} A phase I/II trial in patients with acute myeloid leukemia showed that dosing with cyclosporine before and in combination with daunorubicin prevented chemotherapy resistance, while also resulting in a decrease in MDR-1 RNA expression.²⁴ Our results may highlight one potential treatment approach to MDR tumors that has not yet been fully exploited in the clinical environment, specifically the prevention of the emergence of resistance through the early use of P-gp inhibitors.¹⁻³ It seems reasonable that agents such as dofequidar may be useful in the adjuvant or even neoadjuvant setting with the goal of preventing or delaying the induction of MDR associated with chemotherapy.

The potential clinical significance of P-gp and MRP expression in breast cancer is supported by the results from a number of studies. For example in a study of primary breast cancer patients (n = 259), MRP expression was associated with an increased risk of treatment failure in patients with small tumors (T1) and node-positive patients who received adjuvant cyclophosphamide, methotrexate, and fluorouracil (CMF) chemotherapy but not in node-negative patients.²⁵ Burger et al¹² reported that the expression of MDR1 mRNA in primary breast tumors was inversely correlated with the efficacy of first-line chemotherapy. Additionally, the high level of MDR1 expression was suggested to be a significant predictor of poor prognosis in patients

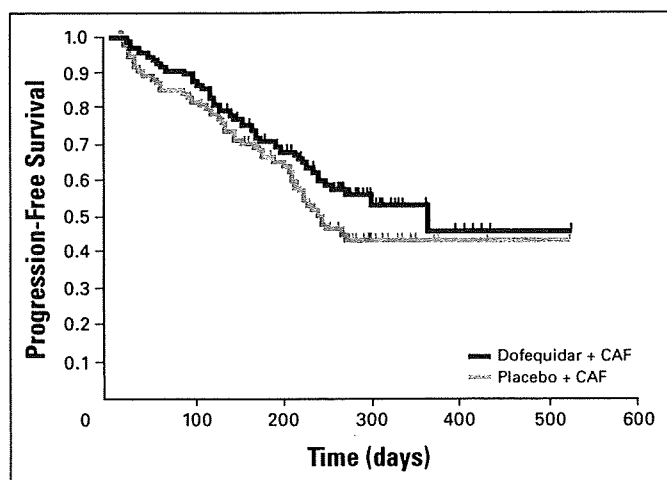


Fig 2. Progression-free survival in patients treated with dofequidar plus cyclophosphamide, doxorubicin, and fluorouracil (CAF) and placebo plus CAF ($P = .145$).

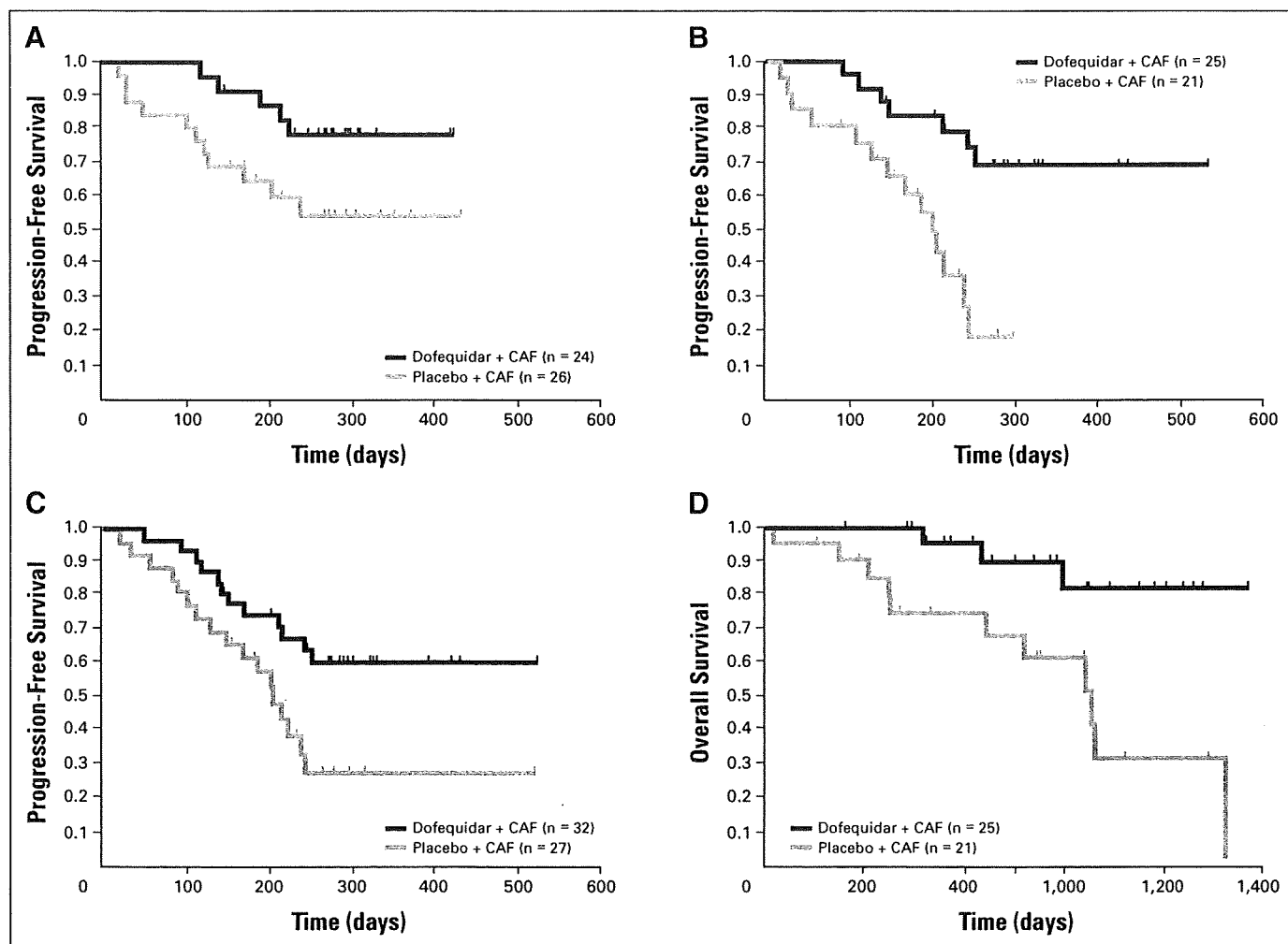


Fig 3. Subgroup analyses. (A) Progression-free survival in premenopausal patients ($P = .046$); (B) progression-free survival in patients who had no prior therapy ($P = .0007$); (C) progression-free survival in patients who were stage IV at diagnosis with an intact primary tumor ($P = .017$); and (D) overall survival in patients who had no prior therapy ($P = .0034$).

with advanced disease.¹² Significantly increased expression of P-gp and MRP-1 has also been reported in an immunohistochemical study of patients treated with preoperative chemotherapy, whereas pretreatment expression of MRP-1 was associated with significantly shorter PFS in patients.²⁶ In a more recent study, MRP-1 expression was shown to be an independent predictor for shorter relapse-free survival and overall survival, after adjuvant CMF treatment, in premenopausal, hormone receptor-positive patients.²⁷ However, MRP-1 expression did not affect patients' response to adjuvant tamoxifen plus goserelin treatment.²⁷

These findings and our results support the view of Leonard et al,³ who indicate that future patients will need to be carefully selected for the identification and development of effective drug-resistance modulators. Patient populations who may derive maximal benefit from MDR inhibition, for example, the no-prior-therapy, advanced-disease, or premenopausal patient group in the present study, could quite easily be overlooked or lost within a large, heterogeneous trial population.³ Furthermore, by refining future clinical trials to incorporate specific disease and patient characteristics, a clearer picture of drug resistance in cancer will be obtained and the most effective MDR inhibitor/chemotherapeutic agent(s) selected.

Many MDR inhibitors have required high serum concentrations for MDR reversal, which resulted in unacceptable toxicity, thereby limiting their clinical impact.^{7,28-32} Although more recent agents have shown improved tolerability profiles, this has been countered by unpredictable pharmacokinetic interactions with other transporter molecules (eg, cytochrome P450-mediated drug metabolism and excretion, necessitating dose reductions in chemotherapy agents and leading to inconsistent chemotherapy dosing among patients).^{1,5} Similarly, the addition of the MDR-modulating agent valspodar (PSC 833) to chemotherapy agents did not improve treatment outcome.^{33,34} Toxicity was increased in the valspodar-treated group compared with chemotherapy agents alone, despite the reduction of chemotherapy doses in the valspodar-containing regimen. In our study, dofequidar was well tolerated, with no indication of the unacceptable toxicity associated with early MDR inhibitors. Importantly, dofequidar did not affect the plasma concentrations of doxorubicin in patients during the study and displayed an acceptable pharmacokinetic profile.

In conclusion, this study suggests that treatment with dofequidar resulted in possible clinical benefit for patients who had not received prior therapy, who were premenopausal, or who were stage IV at diagnosis with an intact primary tumor. Dofequidar was also well

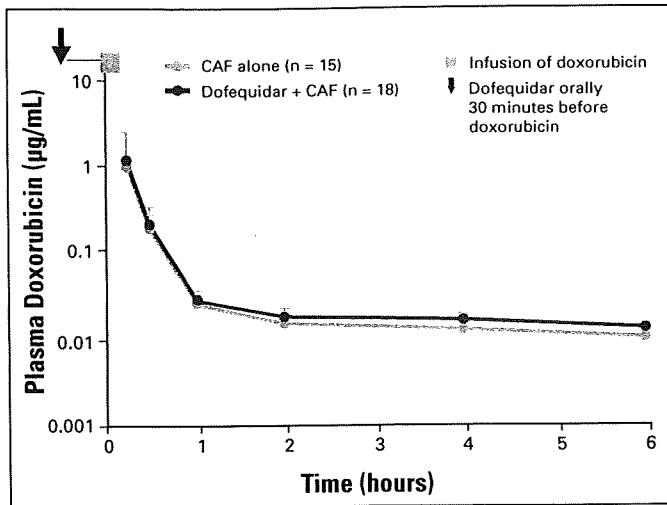


Fig 4. Plasma levels of doxorubicin in patients receiving dofequidar or placebo. CAF, cyclophosphamide, doxorubicin, and fluorouracil.

tolerated in the clinical setting and had no impact on doxorubicin pharmacokinetics. Further studies are merited to assess the effect of dofequidar in specific patient populations with breast cancer.

AUTHORS' DISCLOSURES OF POTENTIAL CONFLICTS OF INTEREST

Although all authors completed the disclosure declaration, the following author or immediate family members indicated a financial interest. No conflict exists for drugs or devices used in a study if they are not being

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REFERENCES

- Gottesman MM, Fojo T, Bates SE: Multidrug resistance in cancer: Role of ATP-dependent transporters. *Nat Rev Cancer* 2:48-58, 2002
- Loe DW, Deeley RG, Cole SPC: Biology of the multidrug resistance-associated protein, MRP. *Eur J Cancer* 32A:945-957, 1996
- Leonard GD, Fojo T, Bates SE: The role of ABC transporters in clinical practice. *Oncologist* 8:411-424, 2003
- Michalak K, Hendrich AB, Wesolowska O, et al: Compounds that modulate multidrug resistance in cancer cells. *Cell Biol Mol Lett* 6:362-368, 2001
- Thomas H, Coley HM: Overcoming multidrug resistance in cancer: An update on the clinical strategy of inhibiting p-glycoprotein. *Cancer Control* 10:159-165, 2003
- Ambudkar SV, Dey S, Hrycyna CA, et al: Biochemical, cellular, and pharmacological aspects of the multidrug transporter. *Annu Rev Pharmacol Toxicol* 39:361-398, 1999
- Krishna R, Mayer LD: Multidrug resistance (MDR) in cancer: Mechanisms, reversal using modulators of MDR and the role of MDR modulators in influencing the pharmacokinetics of anticancer drugs. *Eur J Pharm Sci* 11:265-283, 2000
- Mechetner E, Kyshtobayeva A, Zonis S, et al: Levels of multidrug resistance (MDR1) P-glycoprotein expression by human breast cancer correlate with *in vitro* resistance to taxol and doxorubicin. *Clin Cancer Res* 4:389-398, 1998
- Esteva FJ, Valero V, Pusztai L, et al: Chemotherapy of metastatic breast cancer: What to expect in 2001 and beyond. *Oncologist* 6:133-146, 2001
- Hortobagyi GN: Treatment of breast cancer. *N Engl J Med* 339:974-984, 1998
- National Comprehensive Cancer Network: Clinical Practice Guidelines in Oncology, version 1. Jenkintown, PA, National Comprehensive Cancer Network, 2005
- Burger H, Foekens JA, Look MP, et al: RNA expression of breast cancer resistance protein, lung resistance-related protein, multidrug resistance-associated proteins 1 and 2, and multidrug resistance gene 1 in breast cancer: Correlation with chemotherapeutic response. *Clin Cancer Res* 9:827-836, 2003
- Kroger N, Achterath W, Hegewisch-Becker S, et al: Current options in treatment of anthracycline-resistant breast cancer. *Cancer Treat Rev* 25:279-291, 1999
- Tsuruo T, Iida H, Tsukagoshi S, et al: Overcoming of vincristine resistance in P388 leukemia *in vivo* and *in vitro* through enhanced cytotoxicity of vincristine and vinblastine by verapamil. *Cancer Res* 41:1967-1972, 1981
- Tsuruo T: Circumvention of drug resistance with calcium channel blockers and monoclonal antibodies, in Ozols R (ed): *Drug Resistance in Cancer Therapy*. Norwell, MA, Kluwer Academic Publishers, 1989, pp 73-95
- Tsuruo T, Naito M, Tomida A, et al: Molecular targeting therapy of cancer: Drug resistance, apoptosis and survival signal. *Cancer Sci* 94:15-21, 2003
- Sato W, Fukazawa N, Nakanishi O, et al: Reversal of multidrug resistance by a novel quinoline derivative, MS-209. *Cancer Chemother Pharmacol* 35:271-277, 1995
- Nakanishi O, Baba M, Saito A, et al: Potentiation of the antitumor activity by a novel quinoline compound, MS-209, in multidrug-resistant solid tumor cell lines. *Oncol Res* 9:61-69, 1997
- Narasaki F, Oka M, Fukuda M, et al: A novel quinoline derivative, MS-209, overcomes drug resistance of human lung cancer cells expressing the multidrug resistance-associated protein (MRP) gene. *Cancer Chemother Pharmacol* 40:425-432, 1997
- Margolese RG, Hortobagyi GN, Bucholz TA: Neoplasms of the breast, in Kufe DW, Pollock RE, Weichselbaum RR, et al (eds): *Cancer Medicine*, (ed 6). Hamilton, Canada, BC Decker Inc, 2003
- Wishart GC, Bissett D, Paul J, et al: Quinidine as a resistance modulator of epirubicin in advanced breast cancer: Mature results of a placebo-controlled randomized trial. *J Clin Oncol* 12:1771-1777, 1994
- Belpomme D, Gauthier S, Pujade-Lauraine E, et al: Verapamil increases the survival of patients with anthracycline-resistant metastatic breast carcinoma. *Ann Oncol* 11:1471-1476, 2000
- Cocker HA, Tiffin N, Pritchard-Jones K, et al: *In vitro* prevention of the emergence of multidrug resistance in a pediatric rhabdomyosarcoma cell line. *Clin Cancer Res* 7:3193-3198, 2001
- List AF, Spier C, Greer J, et al: Phase I/II trial of cyclosporine as a chemotherapy-resistance

modifier in acute leukemia. *J Clin Oncol* 11:1652-1660, 1993

25. Nooter K, Brutel de la Riviere G, Look MP, et al: The prognostic significance of expression of the multidrug resistance-associated protein (MRP) in primary breast cancer. *Br J Cancer* 76:486-493, 1997

26. Rudas M, Filipits M, Taucher S, et al: Expression of MRP1, LRP and Pgp in breast carcinoma patients treated with preoperative chemotherapy. *Breast Cancer Res Treat* 81:149-157, 2003

27. Filipits M, Pohl G, Rudas M, et al: Clinical role of multidrug resistance protein 1 expression in chemotherapy resistance in early-stage breast cancer: The Austrian Breast and Colorectal Cancer Study Group. *J Clin Oncol* 23:1161-1168, 2005

28. Bradshaw DM, Arceci RJ: Clinical relevance of transmembrane drug efflux as a mechanism of multidrug resistance. *J Clin Oncol* 16:3674-3690, 1998

29. Ferry DR, Traunecker H, Kerr DJ: Clinical trials of P-glycoprotein reversal in solid tumours. *Eur J Cancer* 32A:1070-1081, 1996

30. Fisher GA, Sikic BI: Clinical studies with modulators of multidrug resistance. *Hematol Oncol Clin North Am* 9:363-382, 1995

31. Fisher GA, Lum BL, Hausdorff J, et al: Pharmacological considerations in the modulation of multidrug resistance. *Eur J Cancer* 32A:1082-1088, 1996

32. Kerr DJ, Graham J, Cummings J, et al: The effect of verapamil on the pharmacokinetics of adria-

mycin. *Cancer Chemother Pharmacol* 18:239-242, 1986

33. Baer MR, George SL, Dodge RK, et al: Phase 3 study of the multidrug resistance modulator PSC-833 in previously untreated patients 60 years of age and older with acute myeloid leukemia: Cancer and Leukemia Group B Study 9720. *Blood* 100:1224-1232, 2002

34. Friedenberg WR, Rue M, Blood EA, et al: Phase III study of PSC-833 (valspodar) in combination with vincristine, doxorubicin, and dexamethasone (valspodar/VAD) versus VAD alone in patients with recurring or refractory multiple myeloma (E1A95): A trial of the Eastern Cooperative Oncology Group. *Cancer* 106:830-838, 2006

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Appendix

The Appendix is included in the full-text version of this article, available online at www.jco.org. It is not included in the PDF version (via Adobe® Reader®).

各論 化学療法

術前化学療法の適応と限界

Indications and limitations of primary systemic therapy
for operable breast cancer

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Key words : 術前化学療法, 手術可能原発性乳癌, primary systemic therapy, operable breast cancer

はじめに

手術可能な原発性乳癌に対する治療戦略は、21世紀に入り大きな転換期を迎えた。早期原発性乳癌に対しては、従来から根治手術後に術後補助化学療法が行われてきたが、化学療法を術前に施行しても、術後に施行しても、無病生存率および全生存率に有意差は認めないという結果が、National Surgical Adjuvant Breast and Bowel Project (NSABP) B-18 トライアル¹⁾および European Organization for Research and Treatment of Cancer (EORTC) 10902 トライアル²⁾から2001年に報告された。また、術前化学療法を行い病理学的完全奏効 (pathological complete response: pCR) が得られた症例は、それ以外の症例と比較して有意に無再発生存期間の延長が認められ、pCRは生存に代わる surrogate endpointとして用いられるようになり、術前化学療法の臨床的有用性が明らかとなった³⁾。このような背景から近年、術前化学療法の重要性を加味してその名称を従来の neoadjuvant/preoperative therapy から primary systemic therapy (PST) とするよう提唱されている。

本稿では、手術可能な原発性乳癌に対する PST をこれまでのエビデンスから考察し、その

適応と限界について言及する。

1. PSTの目的

a. 腫瘍縮小による乳房温存療法の適応拡大

PSTによって原発腫瘍を縮小させ、乳房温存することを目的とする。NSABP B-18 トライアル³⁾では、PST施行群では乳房温存率が67.8%であったのに対し、手術先行群では59.8%で有意に前者の方の温存率が高かった。ただし、乳房内再発率は前者が14.5%、後者が6.9%で、PST後の乳房内再発は約2倍であると報告された¹⁾。しかし、最近ではPST後でも乳房内再発率は高くないという報告もみられる。いずれにしても、PST後の乳房温存療法の適応はMDCTやMRIなどの画像診断を駆使して慎重に決定し、乳房内再発を防止するためには病理組織学的断端陰性を確保することが重要である。

b. 長期無再発生存可能な症例の選別

NSABP B-18¹⁾およびB-27²⁾ トライアルにおけるpCR症例は、それ以外の症例と比較して有意に無再発生存期間の延長が認められた。すなわち、原発巣がpCRであれば全身への微小転移も同時に根絶されたと考えられ良好な長期生存が期待できるため、pCRは生存に代わる surrogate endpointとして用いられるようになった。

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表1 術前化学療法における代表的無作為化臨床試験

トライアル/著者, 年	n	対象	レジメン	ORR (%)	pCR率 (%)	生存率 (%)
NSABP B-18/ Fisher ら ³⁾ , 1998 Wolmark ら ¹⁾ , 2001	1,523	T1-3N0-1M0	AC×4→S vs S→AC×4	79	13.0	69 (9.5年) [#] 70
EORTC 10902/ von der Hage ら ²⁾ , 2001	698	T1c-4bN0-1M0	FEC×4→S vs S→FEC×4	NA	4.0	82 (4.7年) [#] 84
NSABP B-27/ Bear ら, 2003 ⁴⁾ , 2006 ⁵⁾	2,411	T1c-3N0-1M0	AC×4→S vs AC×4→TXT×4→S vs AC×4→S→TXT×4	85.5 91.1 85.5	13.7 26.1 13.7	81 (6.5年) [#] 82
Aberdeen/ Smith ら ⁶⁾ , 2002 Hutcheon ら ⁷⁾ , 2003	162	T2-4N0-2M0 (T>3cm)	CVAP×4→NR→TXT×4 vs CVAP×4→R→TXT×4 vs CVAP×4→R→CVAP×4	47.0 85.0 64.0	1.8 30.8 15.4	NA 97 (5.4年) [#] 78
MD Anderson/ Green ら ⁸⁾ , 2005	258	T1-3N0-1M0	TXLqw×12→FAC×4 vs TXLq3w×4→FAC×4	NA	28.8* 13.6*	NA NA
GEPARUO/ von Minckwitz ら ⁹⁾ , 2005 Raab ら ¹⁰⁾ , 2004	913	T2-3N0-2M0 ≥2cm	AC×4→TXT×4 vs dose dense AT q2w	85 75	14.3* 7.0*	85 (5年) [#] 81

A: doxorubicin, C: cyclophosphamide, E: epirubicin, S: surgery, TXT: docetaxel, V: vincristine, P: prednisolone, NR: no response, R: response, TXL: paclitaxel, F: fluorouracil, NA: not available, *including nodal status, # (median follow up period)

NSABP トライアルおよびその他の代表的無作為化臨床試験を表1に示す。

2. PSTの適応

術後補助化学療法の適応となるすべての症例がPSTの適応になり得る。すなわち、2005年のザンクトガレンのコンセンサスメETINGにおけるリスク分類でintermediate risk以上に入る症例である。臨床的には35歳未満、明らかなリンパ節転移あり、病理学的には腫瘍径(浸潤径)2cm以上、組織学的異型度II以上、高度脈管侵襲およびHER2/neu(HER2)陽性であり、以上のうち1つでも該当するものは適応になり得る。

しかし、現状では臨床的な条件でその適応を決定するのが一般的であり、Stage IIAでも腫瘍径3cm以上の浸潤癌およびStage IIB以上は適応になる。

3. 至適レジメンと至適投与期間

大多数のトライアルで確認されたことは、

アンスラサイクリン(An)系抗癌剤にタキサン(Tx)系抗癌剤を上乗せした方が、pCRを得る割合が高くなることで、おおよそ20%以上のpCRが得られている。特にAn系抗癌剤の効果が認められている場合でも、同じ治療法を継続するよりもTx系抗癌剤に治療法を変更した方がより高い抗腫瘍効果が期待できることがAberdeen トライアル⁶⁾で確認され、非交差耐性薬剤を早期に導入することが重要であると考えられている。

至適投与期間に関しては、様々なトライアルで8-36週の間で計画され、トライアルによっては手術前後に化学療法を施行するように計画されているものもあるが、少なくとも4サイクルは術前に施行すべきである。

4. pCRの定義

a. 原発巣に対する効果判定

欧米では、癌細胞がすべて消失した場合か乳管内病巣のみが残存した場合、すなわち浸潤癌が消失していればpCRと定義していることが

多い。癌細胞が完全に消失した場合 {pCR(all)} と浸潤巣が消失し乳管内病巣のみが残存した場合 {pCR(inv)} の pCR 率は、同一トリアルの中でもかなりの差がある。NSABP B-27 トリアル⁴⁾では、AC(doxorubicin, cyclophosphamide) 4 サイクルのレジメンと AC 4 サイクルに docetaxel 4 サイクルを加えたレジメンの pCR(all) 率はそれぞれ 9.6%, 18.9% であるのに対し、pCR(inv) 率はそれぞれ 13.7%, 26.1% となり、約 1.5 倍 pCR 率が上昇した。トリアル間での pCR 率の比較や、その予後に関する評価も十分注意する必要がある。

b. 腋窩リンパ節 (Ax LN) に対する効果判定

NSABP のトリアルでは原発巣が pCR であれば Ax LN 転移が残存していても pCR と定義しているが、MD Anderson⁹⁾や GEPAR DUO^{9,10)}のトリアルでは原発巣と Ax LN 転移がともに消失した場合を pCR と判定している。NSABP B-18 トリアル³⁾において Ax LN 転移を考慮しない場合の pCR 率は 13% で、考慮した場合は 11% であった。同様に NSABP B-27 トリアル⁴⁾では AC→docetaxel 群における Ax LN を考慮しない場合の pCR 率は 26% であるが、考慮した場合は 22% であった。GEPAR DUO トリアル⁹⁾では、dose dense AT(doxorubicin, docetaxel) と AC→docetaxel の pCR 率を比較し、それぞれ 7% と 14% であったが、Ax LN を考慮しない場合それぞれ 12% と 22% で、Ax LN を pCR の条件に組み入れるか否かで pCR 率に大きな差が出ている。

また、今までのトリアルでは PST 前に Ax LN 転移の有無を確実に評価できていなかったため、Ax LN を pCR の判定に組み込むことにより PST 前から Ax LN 転移がなかった症例も pCR に判定された可能性がある。Hennessy ら¹¹⁾は、5 つの前向き PST 臨床試験において術前穿刺吸引細胞診にて Ax LN 転移が確認された Stage II/III 原発性乳癌 403 症例について Ax LN に対する効果をみたところ、22% の症例に pCR が得られたと報告している。pCR 群と non-pCR 群の 5 年無再発生存率はそれぞれ 87%, 60% で、生存率はそれぞれ 93%, 72% であり、pCR

群で有意に予後良好であった。また、Ax LN の pCR 症例の予後は、原発巣の pCR 達成の有無に影響されなかった。すなわち、PST 後に Ax LN の pCR が達成できれば残存原発巣が認められても予後は良好であることから、原発巣と転移巣には生物学的な違いがあることが示唆された。

5. 腫瘍の生物学的特性による治療効果予測

a. ホルモンレセプター (HR) の有無

大部分のトリアルにおける HR 陰性乳癌に対する PST による pCR 率は、それぞれ陽性乳癌の約 2-4 倍と高く、16-42% であった。また、Nakamura ら¹²⁾は、202 例の手術可能原発性乳癌に対して FEC100 (fluorouracil 500 mg/m², epirubicin 100 mg/m², cyclophosphamide 500 mg/m²) 4 サイクルと docetaxel 75 mg/m² 4 サイクルの順次投与を行い、全体の pCR 率は 23% であったが、HR 陽性かつ HER2 陰性乳癌の pCR 率は 13% と低率であったのに対し、HR 陰性かつ HER2 陽性乳癌の pCR 率は 65% と高率であったと報告している。

b. HER2, topoisomerase II α および Ki-67 の発現

HER2 と topoisomerase II α (Topo II) 遺伝子は 17 番染色体の長腕 (q12-q21) 領域に近接して存在し、HER2 遺伝子の過剰発現した乳癌の 20-30% 程度に Topo II 遺伝子の過剰発現がある。An 系抗癌剤は Topo II を阻害することで抗腫瘍効果をもたらすので、Topo II 遺伝子の過剰発現は、An 系抗癌剤を含む化学療法に対する効果予測因子となると考えられている。PST においても同時増幅例に対しては An 系抗癌剤による pCR 率の向上が期待される。

一方 Petit ら¹³⁾は、免疫染色による Ki-67 の高発現 (20% 以上) は、高い細胞増殖能を反映し An 系抗癌剤を含む PST の効果予測因子として重要であることを報告している。また、Bozzetti ら¹⁴⁾は、An 系抗癌剤の投与量の違い (低用量と高用量)、HER2 遺伝子の過剰発現の有無、HR の有無および免疫染色による Ki-67 の高発現の有無の因子間で PST の臨床効果について

表2 HER2 過剰発現を呈する原発性乳癌に対する trastuzumab 併用術前化学療法

著者, 年	n	対 象	レジメン	cRR(%)	pCR(%)
Burstein ら ¹⁸⁾ , 2003	40	T1-3N0-1M0	12Hqw+4P(175)q3w	75	18
Coudert ら ¹⁹⁾ , 2006	33	T1-3N0-1M0	18Hqw+6D(100)q3w	96	41
Hurley ら ²⁰⁾ , 2006	48	Stage II, III, 炎症性	12Hqw +4{D(70)q3w+Cp(70)}q3w	100	23
佐野ら ²¹⁾ , 2006	21	T>3 cm or N+	12Hqw+4D(75)q3w	90	21
Buzdar ら ²²⁾ , 2005	42*	T1-3N0-1M0	4P(225)q3w→4FEC(75)q3w with or without Hqw	87 vs 47	65 vs 26

P: paclitaxel, D: docetaxel, H: trastuzumab, Cp: cisplatin, FEC: fluorouracil, epirubicin and cyclophosphamide, *randomized, (dose) mg/m²

多変量解析した結果, 免疫染色による Ki-67 の高発現の有無が独立した臨床効果予測因子であったと報告している。

c. Triple negative (TN) 腫瘍

エストロゲンおよびプロゲステロンレセプター (ER および PgR) 陰性, かつ HER2 過剰発現のない乳癌を TN 乳癌と呼称している。ER/PgR と HER2 により定義された腫瘍タイプ間での遺伝子発現プロファイルが異なることは幾つかの報告で明らかとなった。Sorlie ら¹⁵⁾ は, 乳癌のサブタイプを確認するように設計された固有の遺伝子リストを確認し, luminal (管腔), basal-like (基底膜様), HER2 サブタイプなど幾つかの確認可能なクラスターに分類した。更に, 固有の遺伝子リストによって確認された乳癌サブタイプは臨床上的特徴, 転帰および治療に対する反応が異なることが示された。なかでも TN 乳癌のおおよそ 80% は basal-like 腫瘍であり, 予後不良である。これらは内分泌療法や trastuzumab 療法などの乳癌標的治療の対象とならずに化学療法のみが治療手段として残る。PST において, MD Anderson の試験¹⁶⁾ では遺伝子プロファイリングが行われた原発性乳癌 83 例の pCR 率は, luminal 腫瘍より basal-like 腫瘍が有意に高かった。また, UNC 試験¹⁷⁾ では 105 例に対して免疫組織化学的にサブタイプ分類が行われ, luminal, basal-like, HER2 タイプは

それぞれ 52%, 27%, 21% であった。術前 AC 療法を行った結果, pCR 率はそれぞれ 13%, 30%, 27% であり, basal-like 腫瘍で一番高かったと報告している。化学療法に対する感受性を考えると basal-like 腫瘍の予後が不良なのは逆説的にみえるが, UNC 試験¹⁷⁾ における観察期間 2.5 年において basal-like 腫瘍は luminal 腫瘍と比較して無遠隔転移生存率が低く, 全生存率で有意に悪かった。これは, PST に奏効しなかった basal-like 腫瘍は他の化学療法にも反応を示さずに不良な予後をたどることを示唆している。

basal-like 腫瘍に代表される TN 腫瘍に対する PST は, 現在の標準的レジメンである An 系抗癌剤と Tx 系抗癌剤を用いることを基本として今後更に有効なレジメンの開発が必要である。

6. HER2 過剰発現を呈する原発性乳癌に対する trastuzumab 併用 PST

Tx 系抗癌剤と trastuzumab の併用療法で pCR 率が報告されている主な phase II トライアルを表 2 に示す。対象症例は Stage II 以上で, なかには Stage IIIb や炎症性乳癌を対象とした試験もあるためそれぞれの効果の比較は困難であるが, pCR 率は 18-41%¹⁸⁻²¹⁾ と比較的高かった。Tx 系抗癌剤は, 4 サイクル^{18,20,21)} または 6 サイクル¹⁹⁾ 投与され, 6 サイクル投与でより高い pCR

率が得られる傾向があった。本レジメンは、期待される治療法であるが症例数も 20-50 例程度と少数であり、今後予後を含めた多数例での検討が必要である。

Buzdar ら²⁹⁾は、術前に paclitaxel 225 mg/m² を 3 週ごと 4 サイクルのあと FEC75 (fluorouracil 500 mg/m², epirubicin 75 mg/m², cyclophosphamide 500 mg/m²) 3 週ごと 4 サイクル投与するレジメンと、そのレジメンに trastuzumab を毎週 24 回併用したレジメンの 2 群に無作為に分け、化学療法に対する trastuzumab の併用効果を検討した。最終登録数は 42 例で化学療法単独群の pCR 率は 26.3% であるのに対し、trastuzumab 併用群は 65.2% と有意に高率であった。安全性の面で心機能に関しては両群間で差はなかったが、血液学的毒性に関しては Grade 4 の好中球減少が有意に trastuzumab 併用群で多かった。An 系抗癌剤と trastuzumab の併用レジメンの効果は期待されるが、安全性に関してはいまだ確立されたとはいえない。

現在、paclitaxel 毎週投与 12 回に続く FEC 4 サイクル化学療法に最初から trastuzumab を 24 回併用する群と、FEC 4 サイクル後に paclitaxel と trastuzumab を毎週 12 回併用する群で、大規模なランダム化トライアル (NSABP B-41) が行われており、その結果が待たれる。

また、Nakamura ら³⁰⁾が報告したような HR 陰性かつ HER2 陽性乳癌である化学療法に感受性の高い症例に対しては、まず An 系抗癌剤を 4 サイクル行い、その治療効果によって Tx 系抗癌剤に trastuzumab を併用するかどうかを考慮するという治療戦略も考えられる。

7. pCR 例に対する予後予測

NSABP B-27 トライアルにおける Bear ら³¹⁾の

報告では、pCR 症例についてそれぞれ Ax LN 転移個数別 (0 個, 1-3 個, 4-9 個, 10 個以上) に予後を検討した結果、原発巣が pCR でも Ax LN 転移が多いほど予後不良であった。また、非 pCR 症例においても同様の結果であった。すなわち、Ax LN 転移個数は原発巣の pCR とは独立した強力な予後因子であったと述べている。

また、MD Anderson Cancer Center における PST 後に Ax LN も含む pCR を得た原発性乳癌 226 例のレトロスペクティブな多変量解析の検討では、遠隔転移再発に影響を及ぼす独立した因子は Stage IIIB, IIIC および炎症性乳癌、閉経前、Ax LN 郭清個数 10 個以下の 3 つであった²⁹⁾。3 つの独立した予測因子に 1 つも当てはまらない群、1 因子の群、2 因子の群および 3 因子の群に分類すると、10 年無遠隔再発率はそれぞれ 97%, 88%, 77%, 31% で、各群間で有意差を認めた。原発巣および Ax LN で pCR を得た比較的予後良好な症例でも、閉経前の局所進行乳癌症例は遠隔再発に対する注意が必要であろう。

おわりに

手術可能原発性乳癌に対する PST について最近のエビデンスを中心に述べ、考察した。今までと同様に高い pCR 率を目指すレジメンの開発が進められる一方で、腫瘍の生物学的特性に合わせたテーラーメイド医療を実現するための探索が今後更に期待される。また、pCR の有無にかかわらず PST 後の予後予測因子を更に検討し、それに基づいた術後補助療法の適応と治療戦略を探求することが今後の重要な課題である。

■ 文 献

- 1) Wolmark N, et al: Preoperative chemotherapy in patients with operable breast cancer: Nine-year results from National Surgical Adjuvant Breast and Bowel Project B-18. *J Natl Cancer Inst Monogr* 30: 96-102, 2001.
- 2) von der Hage JA, et al: Preoperative chemotherapy in primary operable breast cancer: Results from the European Organization for Research and Treatment of Cancer Trial 10902. *J Clin Oncol* 19: 4224-4237, 2001.

- 3) Fisher B, et al: Effect of preoperative chemotherapy on the outcome of women with operable breast cancer. *J Clin Oncol* 16: 2672-2685, 1998.
- 4) Bear HD, et al: The effect on primary tumor response of adding sequential preoperative docetaxel to preoperative doxorubicin and cyclophosphamide: Preliminary results from National Surgical Adjuvant Breast and Bowel Project Protocol B-27. *J Clin Oncol* 21: 4165-4174, 2003.
- 5) Bear HD, et al: Sequential preoperative or postoperative doxorubicin plus cyclophosphamide for operable breast cancer: National Surgical Adjuvant Breast and Bowel Project Protocol B-27. *J Clin Oncol* 24: 2019-2027, 2006.
- 6) Smith IC, et al: Neoadjuvant chemotherapy in breast cancer significantly enhanced response to docetaxel. *J Clin Oncol* 20: 1456-1466, 2002.
- 7) Hutcheon AW, et al: Docetaxel primary chemotherapy in breast cancer: A five year update of the Aberdeen trial. *San Antonio Breast Cancer Symposium (Abstr #11)*, 2003.
- 8) Green MC, et al: Weekly paclitaxel improves complete remission in operable breast cancer when compared with paclitaxel once every 3 weeks. *J Clin Oncol* 23: 5983-5992, 2005.
- 9) von Minckwitz G, et al: Doxorubicin with cyclophosphamide followed by docetaxel every 21 days compared with doxorubicin and docetaxel every 14 days as preoperative treatment in operable breast cancer: the GEPARUO study of German Breast Group. *J Clin Oncol* 23: 2676-2685, 2005.
- 10) Raab G, et al: Three weekly doxorubicin with cyclophosphamide followed by docetaxel (AC→T) versus 2 weekly doxorubicin and docetaxel (AT) as preoperative treatment in operable breast cancer: First analysis of the event-free survival of the GEPARUO study. *San Antonio Breast Cancer Symposium (Abstr #5047)*, 2004.
- 11) Hennessy BT, et al: Outcome after pathologic complete eradication of cytologically proven breast cancer axillary node metastases following primary chemotherapy. *J Clin Oncol* 23: 9304-9311, 2005.
- 12) Nakamura S, et al: The effect of pathological response of multicenter phase II trial of fluorouracil, epirubicin, cyclophosphamide (FEC100) followed by docetaxel (DOC75) in primary operable breast cancer. *European Breast Cancer Conference (Abstr #370)*, 2006.
- 13) Petit T, et al: Comparative value of tumor grade, hormonal receptor, Ki-67, HER-2 and topoisomerase II alpha status as predictive markers in breast cancer patients treated with neoadjuvant anthracycline-based chemotherapy. *Eur J Cancer* 40: 205-211, 2004.
- 14) Bozzetti C, et al: Evaluation of HER-2/neu amplification and other biological markers as predictors of response to neoadjuvant anthracycline-based chemotherapy in primary breast cancer. *Am J Clin Oncol* 29: 171-177, 2006.
- 15) Sorlie T, et al: Gene expression patterns of breast carcinomas distinguish tumor subclasses with clinical implications. *Proc Natl Acad Sci USA* 98: 10869-10874, 2001.
- 16) Rouzier R, et al: Basal and luminal types of breast cancer defined by gene expression patterns respond differently to neoadjuvant chemotherapy. *San Antonio Breast Cancer Symposium (Abstr #201)*, 2004.
- 17) Carey LA, et al: The triple negative paradox: Primary tumor chemosensitivity of the basal-like breast cancer (BBC) phenotype. *San Antonio Breast Cancer Symposium (Abstr #1023)*, 2004.
- 18) Burstein HJ, et al: Preoperative therapy with trastuzumab and paclitaxel followed by sequential adjuvant doxorubicin/cyclophosphamide for HER2 overexpressing stage II or III breast cancer: A pilot study. *J Clin Oncol* 21: 46-53, 2003.
- 19) Coudert B, et al: Pre-operative systemic (neo-adjuvant) therapy with trastuzumab and docetaxel for HER-2 over expressing stage II or III breast cancer: results of a multicenter phase II trial. *Ann Oncol* 17: 409-414, 2006.
- 20) Hurley J, et al: Docetaxel, cisplatin, and trastuzumab as primary systemic therapy for human epidermal growth factor receptor 2-positive locally advanced breast cancer. *J Clin Oncol* 24(12): 1831-1838, 2006.
- 21) 佐野宗明ほか: HER2過剰発現を呈する進行乳癌に対するDocetaxelとTrastuzumab併用による術前化学療法の検討—JECBC-02 Trial—. *癌と化学療法* 33(10): 1411-1415, 2006.
- 22) Buzdar AU, et al: Significantly higher pathologic complete remission rate after neoadjuvant therapy

- with trastuzumab, paclitaxel, and epirubicin chemotherapy: results of a randomized trial in human epidermal growth factor receptor 2-positive operable breast cancer. *J Clin Oncol* 23: 3676-3685, 2005.
- 23) Gonzales-Angulo AM, et al: Factors predictive of distant metastases in patients with breast cancer who have a pathologic complete response after neoadjuvant chemotherapy. *J Clin Oncol* 23: 7098-7104, 2005.

特集 術前薬物療法のbreak through

JBCRG03:Docetaxel 75mg/m² followed by FEC100mg/m² による術前化学療法

—JBCRG01, 02からのreviewとbreakthrough—

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Here we report on the concept and design of Japan Breast Cancer Research Group (JBCRG)03 trail which resulted from the experience of JBCRG01 ; FEC100 mg/m² followed by docetaxel 75mg/m² and JBCRG02 ; FEC100 mg/m² followed by docetaxel 100mg/m². Our goal is to find the ultimate primary systemic therapy for operable breast cancer. JBCRG01 trial was started in 2002 to evaluate the efficacy and safety of FEC100 followed by docetaxel 75mg/m² for operable primary breast cancer. The subsequent JBCRG02 trial used increasing docetaxel from 75 mg/m² to 100mg/m² to try and improve results obtained in the interim analysis of JBCRG01. Our current study, JBCRG03 was designed as a reverse regimen to resolve some important issues arising from these previous studies. Here we discuss the issues encountered and the rationale for our methodology in this new trial. Further studies will maximize the results obtained in JBCRG01-03.

Key words : Breast cancer, Primary systemic chemotherapy, Clinical trial

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はじめに

乳癌治療において、Fisherの報告を例に挙げるまでもなく一定の条件からは全身性疾患の性格を有し、根治的手術後であっても多くの患者では補助療法が有用である¹⁾。また、メタアナリシスの結果はホルモン療法・化学療法とも再発・死亡のリスクを低下させることを支持している²⁾。

歴史的には、locally-advanceの患者にreduction chemotherapyとして実施してきた術前化学療法は、今や1) *in vivo*での化学療法の感受性試験、2) 腫瘍縮小によるbreast conserving rate (BCR) の向上、3) 病理学的検索による予後のsurrogate markerとして実臨床にも浸透している。われわれはJBCRGで実施したJBCRG01から03の術前化学療法のtrialを通して得られた知見を報告する。

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1. JBCRG01~03の術前化学療法の歴史的背景

1) 世界の術前化学療法の歴史

術前化学療法の歴史においてNational Surgical Adjuvant Breast and Bowel Project (NSABP) の2つのtrialの結果は今日の治療のmilestoneとして世界に大きなインパクトを与えた。1つはNSABP-B18であり、切除可能乳癌(T_{1-3} , N_{0-1} , M_0)1,523例に対し、AC療法(doxorubicin 60mg/m²+cyclophosphamide 600mg/m²) 4サイクルを手術前後に実施した試験である。この試験は1) 手術前後での化学療法実施のタイミングの違いは無再発生存、全生存に差を生じない、2) 術前AC群でpCR(病理学的寛解=癌の完全消失+DCISのみの遺残)が得られた患者では非pCRの患者に比較し無再発生存が有意に良好であるという重要な結果をもたらした³⁾。もう1つはNSABP-B27であり、切除可能乳癌(T_{1c-3} , N_{0-1} , M_0) 2,400例に対し、AC療法4サイクル後にdocetaxel 100mg/m² 4サイクルを術前に投与した場合、pCR(no tumor+DCIS) rateが13.7%から26.1%まで飛躍的に改善することを示し、術前化学療法におけるtaxaneの重要性を位置付けた⁴⁾。

2) 当時のわが国における術前化学療法の現状とJBCRG01の成り立ち

NSABP B-27の結果が発表された2001年当時、多くの乳腺治療医にとって、術前化学療法は手探り的な状態にあったことが同年の癌治療学会のアンケートから伺える。これによるとanthracyclineに関してはAC, EC(epirubicin+cyclophosphamide), FEC(fluorouracil+epirubicin+cyclophosphamide)と種々のレジメンが用いられていたが、心毒性への懸念から効果よりも安全性が重要視され、ACよりもFECが選択される傾向がみられた。しかしepirubicinの汎用doseは20~90mg/m²と様々でglobal trialの結果へ照合し使用する姿勢には乏しかった様である。Taxaneに関してはsequentialではFEC→Taxane(T), AC→T, EC→Tが、concurrentではTAC(docetaxel+doxorubicin+cyclophosphamide)に関心が向けられていた。sequentialとconcurrentの投与方法による毒性の違いやNSABP-B27とTACの結果からsequentialレジメンの浸透がみられようになってきた。これらのtrialの結果や社会事情を受けて、JBCRGではJBCRG01 studyとして切除可能乳癌(T_{1c-3} , N_{0-1} , M_0)に対してFEC100mg/m²×4 followed by docetaxel 75mg/m²×4の安全性と有効性を検討する臨床試験をデザインした(図1)。

3) JBCRG02への発展

世界的に鑑みてdocetaxelのstandard doseは100mg/m²であり、JBCRG 01での75mg/m²の設定は国外のtrialへの整合性が問われていた。また、JBCRG 01での中間解析の病理学的検討において浸潤癌の癌痕組織の中に高度の変化を有する癌細胞がごく少量残存した所見、つまり僅かにpCRに至らない症例が確認されたことから1) anthracyclineとdocetaxelによるsequential chemotherapyでのglobal dose100mg/m²の検証、2) docetaxelの高用量投与によるpCR rateの向上の検討の為にJBCRG02試験としてFEC100mg/m²×4 followed by docetaxel 100mg/m²×4が実施された。この試験にはJBCRG01参加施設の内コンセプトに合意した数施設によって実施された(図1)。

3) JBCRG03のrationaleとstudy design

さらに、JBCRG01の中間解析による結果は、種々の疑問と考察を生み出した。1) FEC先行の治療ではその毒性により引き続きdocetaxelの減量が認められ、忍容性および効果を低下させている可能性があること、2) FEC responderがdocetaxel変更後に増悪した症例が認められたこと、また3) docetaxel先行regimenが海外の試験で発表されており⁵⁾、わが国でのdocetaxel先行の術前化学療法を検討する余地があること、等からJBCRG01のreverse regimenであるdocetaxel 75mg/m²×4 followed by FEC 100mg/m²×4がJBCRG03試験として実施されるに至った(図1)。Primary endpointは病理組織学的効果(pCR rate)、secondary endpointは安全性、臨床効果、乳房温存率、3年無再発生存率であり、予定症例数は130名、予定登録期間は2005年8月から2006年7月までの12カ月であった。

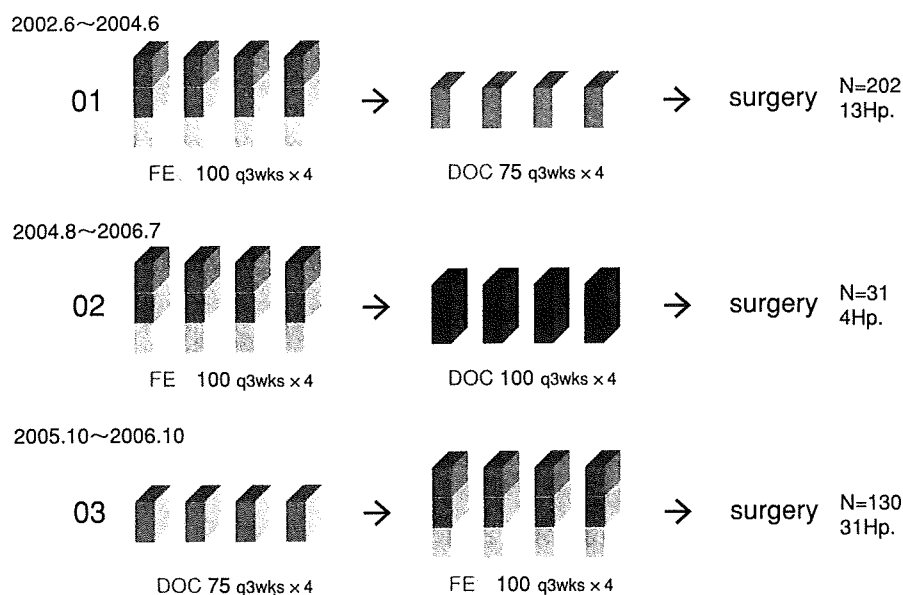


図1 JBCRG neoadjuvant 01-03 study design

2. 結果

JBCRG01およびJBCRG02の結果については、既に報告されているが^{6,7)}、JBCRG03については予定症例数130例に対し、実際の登録期間は2005年10月から2006年10月までの13カ月で137例の登録を得た。今回の検討は137例中111例での解析のため、最終的な効果・認容性に関しては言及しないが、患者背景では閉経前、T2、ERかつ/あるいはPgR陽性が多く登録されていた。血液毒性では白血球・好中球減少はgrade3-4もみられたが、発熱性好中球減少はdocetaxel投与中、FEC投与中いずれも10%前後であった。一方、非血液毒性ではFECにて疲労、悪心・嘔吐および食欲不振においてgrade3が認められたが、他はgrade1-2であり認容性が高かった。全体的に評価するとFECでは血液毒性と消化器症状の発生頻度とgradeが高く、docetaxelでは皮膚症状や浮腫、末梢神経障害などの症状が特徴的であったが、いずれもgradeは低く認容性は高いと考えられた。

3. 考察

1) 01-03から得られた知見

3つの臨床試験から得られた知見をまとめてみる。JBCRG01では当時はわが国においてmanageableか解らなかった世界的標準治療であるFEC100を組み入れ、anthracycline→taxaneでの臨床効果の有用性を検証、JBCRG02ではglobal doseであったdocetaxel 100mg/m²へのdose upによる病理学的効果の向上を検証、JBCRG03ではdocetaxel先行による認容性の向上が臨床的、病理学的効果の向上に寄与するか、reverse regimenとした場合に増悪症例がみられるかが検証されたと言える。

JBCRG01が開始された2002年当時では画期的であったFEC100の実施、pCRを目標とした術前化学療法による臨床試験、中間解析での予想を上回る臨床・病理学的効果等が魅力的であったためか、JBCRG01開始時13施設であったJBCRG参加施設もJBCRG03では31施設まで拡がりをみせた。また、JBCRG01、JBCRG02、JBCRG03の登録期間と症例数を見直すと、JBCRG01では2002年6月から2004年6月までの25カ月で202症例がenrollされたが、JBCRG02試験では24カ月で50症例、JBCRG03では13カ月で130例と症例登録がスムーズとなり、global trialの趨勢を意識した臨床試験参加へのinvestigatorの意欲と術前

化学療法自体の浸透が伺えた。

2) 今後の課題と展望

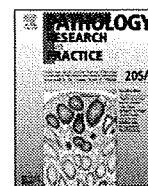
JBCRG01からJBCRG03までの試験の結果は非常に示唆に富むものであった。1) 強力であるFEC100 mg/m²やDocetaxel 75mg/m²を用いても、明らかなchemo resistanceは存在し、この症例における予後の改善が求められる。術前化学療法後に残存病変が有った症例に対する術後治療を検証する試験が存在しないため、われわれはJBCRG04として残存病変がある症例に対するcapecitabine追加投与の有無という試験を計画、始動している。2) JBCRG01試験において治療前に得られた腫瘍の免疫染色の結果から、ER陽性群はpCR rateが低い点が明らかとなった。この結果を受け、特に化学療法の恩恵が乏しいと考えられる閉経患者を対象にaromatase inhibitorの逐次投与の臨床試験が準備されている。3) 同様にJBCRG01試験において諸家が指摘している様にHER2陽性群に高いpCR rateが得られたが、最近特に着目されるtrastuzumabやlapatinib等のHER2 inhibitor, 分子標的療法の追加による更なるpCR rateの向上が期待されている。4) JBCRG01, JBCRG03いずれにおいてdocetaxelのnon-responderが観察されたことから01-03の臨床データとTranslational Research (TR) での種々の情報を組み合わせて、術前化学療法のbest selectionが必須と理解された。

結 語

今回われわれはJBCRGを基盤とした3つの臨床試験から得られた知見を総合的に検証した。種々のコンセプトのもと、3種類のregimenが術前化学療法の試験として遂行され、予定症例数も完了可能であった。慎重な観察のもと、多くが高い忍容性を維持し諸家が報告したpCR rateと同等以上の結果が得られた。これらの試験を通じて、JBCRGは日本での医師主導型臨床試験の普及に寄与したとともに、付随した科学的検証の為にTRを併行するglobal typeの試験の骨子を確立したと言える。

文 献

- 1) Fisher B, Montague E, Redmond C, et al : Comparison of radical mastectomy with alternative treatments for primary breast cancer. A first report of results from a prospective randomized clinical trial. *Cancer* 39 : 2827-2839, 1977
- 2) Early Breast Cancer Trialists' Collaborative Group (EBCTCG) : Effects of chemotherapy and hormonal therapy for early breast cancer on recurrence and 15-year survival : an overview of the randomised trials. *Lancet* 365 (9472) : 14-20, 1687-1717, 2005
- 3) Wolmark N, Wang J, Mamounas E, et al : Preoperative chemotherapy in patients with operable breast cancer : nine-year results from National Surgical Adjuvant Breast and Bowel Project B-18. *J Natl Cancer Inst Monogr* 30 : 96-102, 2001
- 4) Bear HD, Anderson S, Smith RE, et al : The effect on tumor response of adding sequential preoperative docetaxel to preoperative doxorubicin and cyclophosphamide : preliminary results from National Surgical Adjuvant Breast and Bowel Project Protocol B-27. *J Clin Oncol* 15 ; 21 (22) : 4165-4174, 2003
- 5) Gradishar WJ, Wedam SB, Jahanzeb M, et al : Neoadjuvant docetaxel followed by adjuvant doxorubicin and cyclophosphamide in patients with stage III breast cancer. *Ann Oncol* 16 (8) : 1297-1304, 2005
- 6) Nakamura S, Toi M, Takatsuka K, et al : The effect of pathological response of multicenter phase II trial of fluorouracil, epirubicin, cyclophosphamide (FEC100) followed by docetaxel (DOC75) in primary operable breast cancer 5th European Breast Cancer Conference, Nice, 21-25 March 2006
- 7) 濱岡 剛, 中村清吾, 増田慎三, 他 : JBCRG02 : 原発乳がんに対するFEC 100 followed by Docetaxel 100による術前化学療法 2007日本乳癌学会総会プログラム抄録集15回 p247
- 8) Roche H, Fumoleau P, Spielmann, et al : Sequential adjuvant epirubicin-based and docetaxel chemotherapy for node-positive breast cancer patients : the FNCLCC PACS 01 Trial. *J Clin Oncol* 20 ; 24 (36) : 5664-71, 2006



Original Article

Presence of immunoglobulin heavy chain rearrangement in so-called “plasma cell granuloma of the lung”

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SUMMARY

Inflammatory pseudotumor of the lung appears to be a set of heterogeneous disorders. Histologically, three subtypes of pulmonary IPTs have been delineated. Among these, plasma cell granuloma (PCG) is characterized by prominent lymphoplasmacytic infiltration, and PCG has been added to the list of differential diagnostic problems of mucosa-associated lymphoid tissue (MALT) type lymphoma. To investigate the presence or absence of monoclonal B-cell proliferation, we analyzed the immunohistological and genotypic findings in three cases of pulmonary PCGs. Histologically, the three lesions were characterized by severe infiltration of mature plasma cells, plasmacytoid cells, and small lymphocytes intermixed. Scattered Russell bodies (intracytoplasmic inclusions) were present in all three cases, but there were no Dutcher bodies (intranuclear inclusions) or centrocyte-like cells. Immunohistochemical studies of light chain determinants demonstrated the polytypic nature of B-cells. There was no CD5⁺, CD43⁺ or cyclin D1⁺ B-lymphocytes in any of the three lesions. There were no lymphoepithelial lesions detected within any of the three lesions even by immunostaining for cytokeratin. However, polymerase chain assay for immunoglobulin heavy chain gene demonstrated a clonal band in one of the three cases. It currently remains unclear whether this one case, demonstrating IgH gene rearrangement in our series, could be a sign of the prelymphomatous stage (e.g. incipient MALT type lymphoma) or merely represents an exaggeration of normal B-cell clonal response.

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Introduction

So-called “Inflammatory pseudotumors (IPTs)” affect almost all major organs including the lungs [3,20]. Histologically, IPTs are characterized as an irregular proliferation of myofibroblasts intermixed with inflammatory cells, mainly lymphocytes and plasma cells [3,20]. Histologically, IPT of the lung has been classified into three histological types: (i) fibrohistiocytic type, (ii) plasma cell granuloma (PCGs), and (iii) largely sclerosed or fibrosed type [20]. True neoplastic proliferations of mesenchymal or dendritic cells have been reported in some IPTs [2,3,13,21]. Pulmonary IPTs have also been associated with previous viral infections such as human herpes virus type-8 (HHV-8) [8].

Recently, Zen et al. [22] demonstrated that some pulmonary PCGs represent an IgG4-related sclerosing disease. Rarely, pulmonary IPTs have also been associated with B-cell lymphoma [13]. Moreover, PCG has been added to the list of differential diagnostic problems of extramedullary plasmacytoma (EMP) [12]. To examine the presence or absence of monoclonal B-cell proliferation, we analyzed the immunohistological and genotypic findings in three cases of pulmonary PCGs.

Materials and methods

The tissue specimens were fixed in formalin solution, routinely processed and embedded in paraffin. For light microscopic examination, the sections were stained with hematoxylin–eosin (HE) and elastica van Gieson (EVG) stain.

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