Dose-dense epirubicin and paclitaxel with G-CSF: a study of decreasing intervals in metastatic breast cancer

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Summary Anthracyclines and taxanes are very effective drugs in the treatment of advanced breast cancer. With G-CSF support, the dose-intensity of this combination can be increased by reducing the interval between chemotherapy cycles, the so-called ‘shortening of cycle time’. An 8-day interval was not feasible due mainly to incomplete neutrophil recovery at the day of the next scheduled cycle. In the 10-day interval cohort it was feasible to increase the paclitaxel dose to 175 mg m⁻². The haematological and non-haematological toxicity was relatively mild. No cumulative myelosuppression was observed over at least three consecutive cycles. In combination with G-CSF, epirubicin 75 mg m⁻² and paclitaxel 175 mg m⁻² could be safely administered every 10 days over at least three cycles, enabling a dose intensity of 52 and 122 mg m⁻² per week, respectively. © 2000 Cancer Research Campaign

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Anthracyclines and taxanes are amongst the most active single agents in advanced breast cancer. Studies of single agent paclitaxel demonstrate that leucopenia is frequent and dose-limiting in all schedules (3-, 6-, 24-h infusion) evaluated. Randomized trials evaluating the 3- and/or 24-h schedules, each with two paclitaxel doses of 135 and 175 mg m⁻², have revealed that neutropenia is dose- and schedule-dependent, without apparent difference in efficacy (Eisenhauer et al, 1994; Nabholtz et al, 1996; Gianni et al, 1995a). The optimal therapeutic schedule of paclitaxel is still unknown and, on practical grounds, the short infusion over 3 h is safe, convenient and effective (Gianni, 1999b).

Paclitaxel at longer infusion rate (≥ 24 h) has been given in combination with doxorubicin. The maximal tolerable dose (MTD) and toxicity profile (primarily mucositis and neutropenia) of this combination appear to depend on the sequence and infusion duration of the two drugs (Fisherman et al, 1993; Sledge Jr et al, 1994). In a study of escalating doses paclitaxel (3-h infusion) in combination with fixed dose of doxorubicin (60 mg m⁻² i.v. bolus) in a 3-weekly schedule, mucositis, long-lasting grade 4 neutropenia and febrile neutropenia defined the MTD of paclitaxel at 200 mg m⁻², and was not sequence dependent. This regimen was very effective in chemotherapy-naive breast cancer patients, but at the cost of considerable increased cardiotoxicity (Gianni et al, 1995c; Gehl et al, 1996). 4'-Epi-doxorubicin (epirubicin) is a synthetic doxorubicin analogue, with similar anti-tumour activity as doxorubicin at equimolar doses, but decreased overall toxicity, in particular cardiotoxicity (Brambilla et al, 1986; Jain et al, 1985). It may, therefore, be an attractive substitute for doxorubicin in combination with taxanes.

Granulocyte colony stimulating factor (G-CSF) can be used to ameliorate neutropenia and to obtain increased dose-intensity by allowing a higher dose of chemotherapy per cycle (dose-escalation) or by allowing a shortening of the interval between cycles (dose-dense). Both approaches may lead to a higher dose-intensity, but their biological effect and clinical relevance may be quite different (Henderson et al, 1988). Dose-dense chemotherapy may be important to overcome the cellular cytokinetic resistance of tumors (Gilewski and Norton, 1996). Dose-escalation is based on a steep dose–response relationship, whereby the doses of chemotherapy are increased up to the limits of haematological and non-haematological toxicity. These two approaches were investigated by our group in the treatment of metastatic breast cancer with epirubicin and cyclophosphamide. With the addition of G-CSF, interval reduction permitted a higher dose-intensity, with less toxicity, than dose escalation (Lalisang et al, 1997). Based on these results we started a study with the aim of increasing the dose-intensity of a conventional dose of epirubicin (75 mg m⁻² i.v. bolus) and paclitaxel (135 mg m⁻² 3-h infusion) combination by shortening of the cycle time. We intended to define the minimal tolerable interval of this chemotherapy regimen in combination with G-CSF support. During the study the protocol was amended, and allowed testing a higher paclitaxel dose of 175 mg m⁻² in the shortest feasible interval. The second aim was to assess the safety profile of this approach.
PATIENTS AND METHODS

Patient selection

The study protocol was approved by the institutional review boards of the participating hospitals, and patients gave informed consent. Women with advanced breast cancer had to fulfil the following inclusion criteria:

- Histological proof of breast cancer
- Age 18–70 years
- Performance status 0–2 (Eastern Cooperative Oncology Group scale)
- Neutrophil count \( \geq 2 \times 10^9 \text{l}^{-1} \) and platelet count \( \geq 100 \times 10^9 \text{l}^{-1} \)
- Adequate function tests for liver (bilirubin level \( \leq 25 \mu\text{mol l}^{-1} \) and transaminase level \( \leq 3 \times \text{upper limit of normal} \)) and kidneys (serum creatinine level \( < 150 \mu\text{mol l}^{-1} \))
- No prior chemotherapy for metastatic disease
- Prior adjuvant chemotherapy allowed, if interval last chemotherapy cycle \( \geq 1 \) year and at entry lifetime cumulative dose of doxorubicin \( \leq 300 \text{mg m}^{-2} \) and epirubicin \( \leq 450 \text{mg m}^{-2} \)
- Prior radiotherapy involving \( \leq 25\% \) of red bone marrow
- Left ventricle ejection fraction (LVEF) by multigated (MUGA) isotope cardiograph \( \geq 50\% \) and without symptomatic cardiovascular disease
- No central nervous system involvement.

Treatment plan

In this schedule-finding study, the epirubicin (Pharmacia & Upjohn, Milan, Italy), paclitaxel (Taxol, Bristol-Myers Squibb Pharmaceuticals, Princeton, NJ, USA) as well as R-metHuG-CSF (Filgrastim, Amgen Inc., Thousands Oaks, CA, USA) doses were kept constant, and four intercyclic intervals were foreseen. The starting interval of 14 days was planned to be decreased to 12, 10 and 8 days, respectively. To prevent hypersensitivity reactions due to paclitaxel, a routine premedication regimen was adopted: oral or intravenous dexamethasone \( 20 \text{mg} (6 \text{ and } 12 \text{ h pre-treatment}) \), clemastine \( 2 \text{ mg} \) and ranitidine \( 50 \text{ mg} \) both intravenously 30–60 min before paclitaxel administration. Epirubicin was given as a short i.v. infusion on day 1 at a fixed dose of 75 \( \text{mg m}^{-2} \), Paclitaxel at a dose of 135 \( \text{mg m}^{-2} \) was administered by a 3-h infusion, starting 5 min after epirubicin administration. G-CSF (300 \( \mu\text{g} \) for patients \( \leq 70 \text{ kg} \) and 480 \( \mu\text{g} \) for patients \( > 70 \text{ kg} \)) was administered once daily subcutaneously on all days except the days of chemotherapy.

A cohort of at least six patients was studied at each interval. The neutrophil \( (\geq 2 \times 10^9 \text{l}^{-1}) \) and platelet \( (\geq 100 \times 10^9 \text{l}^{-1}) \) counts had to have recovered on the day of scheduled chemotherapy. For this study specific dose-intensity limiting criteria (DILCs, see Table 1) had been defined. Proceeding to the next, shorter, interval level was only done after completion of the previous cohort and if less than 50\% of those patients had experienced a DILC during the first three courses. In case of incomplete haematological recovery treatment was delayed. Concomitant hormonal therapy or prophylactic antibiotic therapy was not allowed. Patients were transfused when necessary to maintain a platelet count of \( \geq 15 \times 10^9 \text{l}^{-1} \) and haemoglobin level \( \geq 6 \text{ mmol l}^{-1} \).

The patients had to complete a minimum of three cycles, except if one of the following events occurred: disease progression, DILC, any other unacceptable toxicity precluding further therapy, or patients’ refusal to continue treatment. After completion of the first three cycles further therapy was left to the discretion of the investigator.

As the 8-day interval appeared to be not feasible (see Results) a new cohort was tested with a 10-day interval but using a higher paclitaxel dose of \( 175 \text{mg m}^{-2} \).

The minimal tolerable interval (MTI) for this study was defined as the shortest interval that resulted in less than 50\% instances of DILCs among treated patients in each cohort. To express the dose-intensity of the treatment the equation, delivered dose-intensity (DDI), the actually given dose per \( \text{m}^2 \) per week during the first three protocol cycles of treatment, was used.

Pre-treatment and follow-up evaluation

All patients were initially evaluated with a history, physical examination, complete blood-cell count (CBC), liver and kidney function tests, ECG, LVEF MUGA scan, chest X-ray and bone scan. If indicated additional radiological examinations of suspected areas, with tumour measurements (if possible) were performed. CBC was repeated twice weekly and a biochemical profile was assessed before each cycle. Follow-up LVEF MUGA scan was requested after three cycles, in case of clinical signs of congestive heart failure, if patients went off study or at a cumulative dose of 500 \( \text{mg m}^{-1} \), at 800 \( \text{mg m}^{-2} \) of epirubicin and subsequently before each additional treatment course thereafter. In our study cardiac toxicity was defined as development of clinical cardiac failure and/or an absolute decrease in MUGA LVEF \( \geq 20\% \) (EF absolute units) from baseline to a value above 50\% or \( \geq 10\% \) (EF absolute units) to a value below 50\%.

RESULTS

Patient characteristics

Forty-one eligible patients were entered, and five patients were found to be not evaluable; four patients in the 8-day interval

| Table 1 Dose-intensity limiting criteria |
|----------------------------------------|
| 1 Any WHO grade 3 or 4 non-haematological toxicity |
| 2 Neutropenia grade 4: neutrophil count \( < 0.5 \times 10^9 \text{l}^{-1} \) for a period of more than 7 days |
| 3 Febrile neutropenia: neutrophil count \( < 0.5 \times 10^9 \text{l}^{-1} \) and fever |
| 4 Thrombocytopenia grade 4: platelet count \( < 25 \times 10^9 \text{l}^{-1} \) for more than 4 days |
| 5 Delay of chemotherapy due to incomplete recovery on the day of scheduled therapy: Haematological: neutrophil count \( < 2 \times 10^9 \text{l}^{-1} \) and/or platelet count \( < 100 \times 10^9 \text{l}^{-1} \) Persistence of non-haematological side-effects of WHO grade 2 or more (excluding alopecia and anticipatory nausea and vomiting) |
| 6 Cardiotoxicity, defined by development of clinical cardiac failure or an absolute decrease in MUGA LVEF \( \geq 20\% \) (EF absolute units) from baseline to a value above 50\% or \( \geq 10\% \) (EF absolute units) to a value below 50\% |
Cumulative toxicity

In the 14-day interval all five patients without a DILC in the first three cycles continued at their scheduled interval for at least six cycles. In the 12-day interval seven out of eight patients continued the same chemotherapy combination after three protocol cycles, although only two patients at scheduled interval for a total of four cycles, all without additional toxicities.

In the 10-day interval three out of 10 patients continued scheduled treatment for a total of four (two patients) and nine (one patient) cycles, respectively. The single patient with an incomplete neutrophil recovery after cycle three developed a reversible neuropathy grade 3 after cycle 6. In the 8-day interval the two patients without a DILC in the initial three cycles continued the scheduled interval for a total of five and six cycles, respectively. The patient with a febrile neutropenia and stomatitis grade 3 after the second cycle also developed pulmonary embolism. In the 10-day interval at 75/175 mg m\(^{-2}\) only one patient continued scheduled treatment for six cycles. The kinetics of neutrophil and platelet counts for the patients in the various intervals are shown in Figure 1. The non-haematological toxicities were generally mild and are displayed in Table 4. Only one patient encountered a hypersensitivity reaction WHO grade 2.

Transfusion of blood products

In this dose-dense schedule, during the first three cycles red blood-cell (RBC) transfusions were needed in five patients. Most transfusions occurred in the patients receiving more than three cycles (14 of 27 patients). Platelet transfusions were not given.

Cardiotoxicity

A total of 98 MUGA scans were performed; basal and follow-up scans were available in 35 of 36 patients, making them evaluable for cardiotoxicity by ejection fraction. A pathological decline in LVEF, as defined earlier, was observed in two (6%) patients in the first three cycles, both not pre-treated with anthracyclines. The first patient (previously irradiated parasternally) was treated in the 14-day interval, developed a delayed platelet recovery after the first cycle, received cycle 2 after platelet recovery, which was complicated by severe dyspnuea and anaemia (4.5 nmol l\(^{-1}\)), with a transient decrease in LVEF (63\(\rightarrow\)49%). On physical examination, chest X-ray and ECG no signs of congestive heart failure were observed. After RBC transfusion the patient recovered and a control LVEF showed normalization (62%). This patient continued scheduled treatment for an additional three cycles, without further signs of cardiac failure. A short interval (1 day) between irradiation of the lumbar spine and start of study treat-
ment may explain these transient DILCs. In the second patient (previously irradiated to the right chest wall) after three cycles in the 10-day interval at a dose of 75/175 mg m⁻² an asymptomatic decrease in LVEF (60 → 49%) was observed. The patient continued with three cycles of epirubicin/cyclophosphamide with complete normalization of the LVEF. From 20 patients cardiac evaluation was available with a cumulative epirubicin dose of ≥ 450 mg m⁻²; median baseline LVEF 60% (range 50–72%) to 58% (range 30–70%). One patient in the 14-day interval (not previously irradiated) developed congestive heart failure with an LVEF of 30% after cycle 13 (cumulative epirubicin dose 975 mg m⁻²). The single patient with a DILC in the 10-day interval at a dose of 75/135 mg m⁻² (previously irradiated to right chest wall and parasternal region) developed symptomatic ventricular extrasystoles after cycle 3, without a decrease in LVEF, and continued treatment for a total of six cycles. There was no real difference in median changes of LVEF from baseline and after chemotherapy between patients previously irradiated to the loco-regional breast/chest (27 patients, −4%, range –3– +16) and not irradiated patients (8 patients, –7%, range –24 – +9).

Response rate

Although efficacy was not an aim of this study, formal UICC criteria were applied to the 29 patients with measurable disease. After three cycles, i.e. evaluation 20–42 days after the start of the study treatment an objective response (all partial) was already observed in 17 of 29 patients (59%, 95% CI, 41–77%), and there was one patient with progressive disease. Seventeen patients (stable disease 11 and partial response six) continued protocol treatment with an interval of ≤ 14 days for a median number of six cycles (range 4–9) whereby six additional patients reached a partial response and two partial responders reached a complete remission. The median interval between the start of therapy and the first observation of an objective response was 5 weeks (range 3–12 weeks). The best objective response (complete and partial) for the total study group was 79% (95% CI, 65–94%).

DISCUSSION

The aim of our study was to determine the maximal dose intensity of epirubicin in combination with paclitaxel in a dose-dense schedule, supported by G-CSF. With a regimen consisting of epirubicin 75 mg m⁻² and paclitaxel 135 mg m⁻² an interval of 10 days was feasible, allowing a median DDI of 52 mg m⁻² per week for epirubicin and 94 mg m⁻² per week for paclitaxel, respectively. Within the 10-day interval it was feasible to increase the paclitaxel dose to 175 mg m⁻² enabling a median DDI of 122 mg m⁻² per week for paclitaxel in combination with 52 mg m⁻² per week of epirubicin. The treatment-schedule related toxicity was relatively mild, considering the high dose-intensity achieved.

The MTDs of the epirubicin and paclitaxel (3 h) combination in a 21-day schedule without ‘prophylactic’ haematopoietic growth factor as first-line treatment in metastatic breast cancer has been defined at 50/250 mg m⁻², 60/175 mg m⁻², 90/175 mg m⁻² and 90/200 mg m⁻², respectively. Dose-limiting toxicities were primarily haematological, namely severe neutropenia and febrile neutropenia (Catimel et al, 1996; Luck et al, 1997; Conte et al, 1997). The DDI in the latter study is 30 mg m⁻² per week for epirubicin and 67 mg m⁻² per week for paclitaxel, respectively. Dose-limiting toxicities were primarily haematological, namely severe neutropenia and febrile neutropenia (Catimel et al, 1996; Luck et al, 1997; Conte et al, 1997). The DDI in the latter study was 30 mg m⁻² per week for epirubicin and 67 mg m⁻² per week for paclitaxel, which is considerable lower than our results of 52 and 122 mg m⁻² per week, respectively.

Table 4 Number of patients (percentage) encountering non-haematological toxicities (WHO-grading) for all cycles with epirubicin and paclitaxel with G-CSF support

| WHO grading | Nausea/vomiting | Diarrhoea | Stomatitis | Neurotoxicity | Myalgia | Infection | Skin |
|-------------|-----------------|-----------|------------|---------------|---------|-----------|------|
| 1           | 12 (32%)        | 3 (8%)    | 8 (22%)    | 9 (24%)       | 10 (27%)| 0         | 0    |
| 2           | 12 (32%)        | 5 (14%)   | 8 (22%)    | 3 (8%)        | 7 (19%) | 2 (5%)    | 3 (8%)|
| 3           | 1 (3%)          | 0         | 3 (8%)     | 1 (3%)        | 1 (3%)  | 2 (5%)    | 0    |
| 4           | 0               | 0         | 0          | 0             | 0       | 0         | 0    |

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In a study of the epirubicin-paclitaxel combination in a 21-day interval without G-CSF the neutrophil-nadir occurs at day 12 and the median neutropenia grade 4 duration was 3 days (Conte et al., 1996). In our study, with G-CSF, the neutrophil-nadir was reached earlier, at day 8, was of very short duration, and was less pronounced in each consecutive cycle. G-CSF administered on all days, with the exception of the day of chemotherapy, hastened the neutrophil recovery after each chemotherapy course, but may also have had a pre-emptive effect on each consecutive cycle by expansion of the progenitor pool. This so-called pre-emptive G-CSF effect was, however, not observed earlier (Tjan-Heijnen et al., 1998; de Wit et al., 1996). Thrombocytopenia has not been dose-limiting. An important aspect of studies on escalated dose intensities is whether this can be maintained over repeated cycles. Consecutively with intervals as short as 10 days and retreatment at the moment of rapid recovery, the population of progenitor cells may be vulnerable to the repetitive cytotoxic insults and may become exhausted after a number of cycles. Although most data were collected over three cycles, we have not observed signs of exhaustion, even in the small number of patients treated up to six cycles. It may well be that treatment with G-CSF up to the day before the next chemotherapy has protected progenitor cells by putting them out of the cell cycle, as has been observed with GM-CSF (Kobrinsky et al., 1999; Vadhan-Raj et al., 1992). Further data on larger numbers of patients with even more cycles are needed to determine the absence of cumulative myelosuppression with certainty.

In the three epirubicin/paclitaxel studies the objective response rates after a least six cycles were 44%+, 68% (CR in 17%) and 84% (CR in 18%), respectively (Catimel et al., 1996; Luck et al., 1997; Conte et al., 1997). These data suggest a dose–response relationship for epirubicin. In our study, after three short interval cycles a response rate of 59% had already been observed.

An important argument to investigate epirubicin in combination with paclitaxel is to circumvent the increased cardiotoxicity of the doxorubicin/paclitaxel combination. After a median cumulative doxorubicin dose of 480 mg m⁻², 50% of the patients had reductions of the LVEF below the norm and 20% of the patients developed a dense drug delivery. The efficacy and clinical relevance of this approach is now being investigated in a phase II study.

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