Adjuvant Dose-Dense Chemotherapy in Breast Cancer: Standard of Care in High-Risk Patients

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Introduction

Dose-dense chemotherapy plays a controversial role in the adjuvant treatment of breast cancer patients. Whereas meta-analyses [1, 2] persistently describe a significant superiority for dose-dense treatment, the results of large phase III trials remain contradictory [3–9]. Some of these trials showed important differences between the dose-dense and conventional groups regarding number of cycles, type of drug, and total dose. Other trials are accepted and interpreted as dose-dense but present a mixture of dose-dense and conventional schedules [7, 8]. Furthermore, dose-dense and intense dose-dense concepts should be differentiated (fig. 1). Follow-up duration varies between 3 and 10 years, and the risk profiles of the included patient populations can be remarkably different, thus rendering interpretation of the observed results difficult.

Two very relevant points should be stressed before discussing the trials in detail:

Due to the obligatory use of granulocyte colony-stimulating factor (G-CSF) or pegfilgrastim, treatment-related mortality in all trials is consistently lower in dose-dense regimens in comparison to standard schedules. The rate of myelodysplastic syndrome (MDS) or secondary acute myeloid leukemia (AML) is low and corresponds to standard regimens [3, 6, 10]. Provided that the same regimen is given in the dose-dense and the standard arm, there appears to be no evidence that G-CSF or pegfilgrastim are linked to an elevated risk of MDS or AML [3, 4].

Keywords
Dose-dense · Adjuvant chemotherapy · Breast cancer

Summary
Meta-analyses persistently confirm the superiority of dose-dense chemotherapy in comparison with standard chemotherapy. In contrast, individual studies have shown conflicting results. These may be explained by different risk profiles of the treated patient populations. Some trials show a significant advantage in disease-free survival (DFS) and overall survival (OS) in the estrogen receptor (ER)-negative population only, whereas trials with high-risk populations like GIM-2 (Gruppo Italiano Mammella) and AGO-iddETC (Arbeitsgemeinschaft Gynäkologische Onkologie, intense dose-dense epirubicin, paclitaxel, and cyclophosphamide) show a significant superiority in DFS and OS for both, ER-negative and ER-positive patients even after 7 and 10 years, respectively, of follow-up. In contrast, the 10-year follow-up data of the E1199/Intergroup trial no longer showed any superiority of weekly paclitaxel for ER-positive/HER2-negative patients; superiority was observed in the triple-negative subgroup only. Although a direct head-to-head comparison is missing, iddETC or 4 cycles each of dose-dense epirubicin/cyclophosphamide followed by paclitaxel are the preferred adjuvant regimens for patients at risk. Patients with \( \geq 4 \) positive lymph nodes should preferentially be treated with iddETC.

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Adjuvant Dose-Dense Therapy with G-CSF or Pegfilgrastim Support versus Conventional-Dose Chemotherapy

Pure Dose-Dense Concepts versus Conventional-Dose Chemotherapy

Four large randomized trials compared adjuvant dose-dense versus conventionally dosed chemotherapy [3, 4, 9, 11]. In 2 fur-
the trials, the so-called ‘dose-dense’ arm was a mixture of dose-dense and standard-dose chemotherapy, thus hindering an explicit interpretation of the results [7, 8].

The first trial, which showed superiority of dose-dense over conventional-dose chemotherapy, was the Cancer and Leukemia Group B (CALGB) trial 9741 [3] demonstrating that a change in the interval of anthracycline- and taxane-based chemotherapy from every 3 weeks (q3w) to every 2 weeks (q2w) significantly improved disease-free survival (DFS) (hazard ratio (HR) 0.74; p = 0.010) and overall survival (OS) (HR 0.69; p = 0.013) in women with lymph node-positive early-stage breast cancer.

The findings of the recently published GIM-II trial [4] from Italy support the results of CALGB 9741. 2,091 patients were randomized to 1 of 4 study groups: 4 cycles of epirubicin (E)/cyclophosphamide (C) (90/600 mg/m²), followed by 4 cycles of paclitaxel (P) 175 mg/m², given either at 2-week (dose-dense) or 3-week intervals (conventional-dose) plus/minus 5-fluorouracil (5-FU) in addition to EC (FEC). The addition of 5-FU had no impact on DFS and OS, but 5-year DFS (HR 0.77; p = 0.004) and OS rates (HR 0.65; p = 0.001) were significantly better in the dose-dense group. 5-year OS was 94% in the dose-dense arm versus 89% in the standard arm. However, both trials had a standard arm (4 cycles of doxorubicin/cyclophosphamide (AC)/EC followed by 4 cycles of paclitaxel q3w) which probably no longer reflects today’s standard of care.

Four trials did not show superiority of the dose-dense schedule over conventional-dose chemotherapy [7–9, 11]. At first glance, this constellation does not appear to support the superiority of dose-dense chemotherapy. However, can these conflicting results be explained?

The GONO-MIG (Gruppo Oncologico Nord Ovest-Mammella InterGruppo) trial [11] compared FEC q3w versus the same regimen given q2w. The difference between these 2 arms did not reach statistical significance for either recurrence (HR 0.88; p = 0.22) or death (HR 0.87; p = 0.29). These negative results may be explained by the study design. The GONO-MIG trial applied a substandard version of the FEC regimen (i.e., 5-FU at 600 mg/m², epirubicin at 60 mg/m², and cyclophosphamide at 600 mg/m² for 6 cycles). The total dose of epirubicin in both arms was 360 mg/m², which corresponded to only 50% of the total dose in the highly effective Canadian CEF regimen [10].

Results from the UK TACT2 (United Kingdom Trial of Accelerated ChemoTherapy) trial have so far been published in abstract form only [7], and the regimens used in the Canadian MA-21 trial [8] were only in part dose-dense. The TACT2 trial compared 4 cycles of epirubicin q3w versus q2w, followed by standard chemotherapy with either CMF (cyclophosphamide, methotrexate, 5-FU) or capecitabine, and showed no difference in 5-year event-free survival (HR 0.96; p = 0.60) or OS (HR 1.13; p = 0.23).

The 3-arm Canadian MA-21 trial [8] had only a short follow-up duration of 30 months. Patients in the intense ‘dose-dense’ arm received 6 cycles of epirubicin 120 mg/m² and cyclophosphamide 830 mg/m² q2w followed by 4 standard courses of paclitaxel 175 mg/m² q3w (ECQT). Similar to the UK TACT2 trial, the ‘dose-dense’ arm again represents a mixture of dose-dense and standard schedules. The ‘dose-dense’ arm was randomized against 2 standard regimens, i.e. Canadian CEF or AC ×4 followed by paclitaxel ×4 (ACQT). The results of the Canadian trial showed that ACQT was inferior to both CEF and intense dose-dense ECQT. The authors reported no difference in DFS between CEF and intense dose-dense ECQT (HR 0.89; p = 0.46). There are several possible explanations for these results. The total planned cumulative dose, dose intensity, and duration of anthracyline treatment were greater in ECQT and CEF in comparison to ACQT. In the ECQT and ACQT arms, the taxane dose and schedule were the same. Again, the total planned cumulative dose and the dose density were greater in ECQT.

The NSABP B-38 trial [9] was a 3-arm study which compared 4 cycles of dose-dense doxorubicin and cyclophosphamide followed by 4 cycles of dose-dense paclitaxel (dose-dense ACÓP) or the same regimen with 4 cycles of gemcitabine (G) added to paclitaxel (dose-dense ACÓPG) or 6 cycles of docetaxel, doxorubicin, and cyclophosphamide (TAC) q3w. In this large study with nearly 5,000 randomized patients, the addition of gemcitabine to dose-dense ACÓP did not improve outcomes. In addition, no significant differences in efficacy end points were identified between the investigational arm dose-dense ACÓPG and TAC (5-year DFS: HR 0.93; p = 0.39; 5-year OS: HR 0.86; p = 0.17).

Intense Dose-Dense Concepts versus Conventional-Dose Chemotherapy

The importance of dose intensity for adjuvant chemotherapy in patients with breast cancer was first described by Hryniuk as early as 1986 [12]. Higher dose intensity can be achieved by either increasing the single dose per cycle (i.e., higher dose) and/or by reducing the intervals between cycles (i.e., higher dose density).

In contrast to CALGB 9741 and GIM II, which were purely based on the concepts of dose density and sequential application with the total dose of each agents identical in both arms, the intense dose-dense regimen of epirubicin 150 mg/m² q2w ×3, paclitaxel 225 mg/m² q2w ×3, and cyclophosphamide 2,500 mg/m² q2w ×3 (iddETC) was dose-dense and used a higher total dose per cycle [6]. The control arm with 4 cycles EC (AC) followed by 4 cycles of paclitaxel is identical to CALGB 9741 and GIM II.
The iddETC trial recruited only high-risk patients with ≥ 4 positive lymph nodes and is the only trial which reports long-term survival data [5]. With 10 years of follow-up, event-free survival was significantly longer for the iddETC arm with 56 vs. 47% and a HR of 0.74. The relative risk of mortality was reduced by 28% (HR 0.72; log rank test p = 0.0007). Given the fact that 42% of the patients had ≥ 10 positive nodes with a median number of 8 positive nodes, the 10-year survival rate of 69% in the iddETC arm represents to the best of our knowledge the highest survival rate reported for such a high-risk group of patients to date.

Comparison of (Intense) Dose-Dense versus Conventional-Dose Chemotherapy – Why Do We Have Positive and Negative Trials?

A central reason for the conflicting results reported by trials comparing (intense) dose-dense versus standard regimens is likely to be the differences in the risk profile of the studied patient populations (fig. 2). CALGB 9741, GIM-2, and iddETC have shown superiority of (intense) dose-dense regimens. In contrast, UK TACT 2 and NSABP B-38 failed to show superiority of dose-dense regimens. We cannot exclude that the TAC regimen used in the NSABP B-38 trial may be a more effective standard regimen in comparison to AC (EC) followed by paclitaxel which was the standard regimen in the CALGB 9741, GIM-2, and AGO-iddETC trials. A second probable explanation for the contradictory results is the completely different risk profile of the patient populations in the negative and the positive trials.

Whereas the UK TACT 2 trial consists to 87.3% of patients with node-negative or N1 disease, none of these low-risk patients have been recruited by the AGO-iddETC trial. The median number of positive nodes is 1 in the UK TACT 2 and 8 in AGO-iddETC. We cannot exclude that the TAC regimen used in the NSABP B-38 trial may be a more effective standard regimen in comparison to AC (EC) followed by paclitaxel which was the standard regimen in the CALGB 9741, GIM-2, and AGO-iddETC trials. A second probable explanation for the contradictory results is the completely different risk profile of the patient populations in the negative and the positive trials.

Fig. 2. Risk profile of dose-dense trials.

This observation is valid even inside a trial population. In the iddETC trial, patients with ≥ 10 positive lymph nodes showed a higher reduction in mortality compared to patients with 4–9 positive nodes. The same was noted in the GIM-2 trial. Only patients with 4 and more positive nodes had a significant reduction in DFS and OS with dose-dense treatment.

We conclude that the predominant part of the patients recruited by TACT 2 and NSABP B-38 belong to the low- or intermediate-risk group. With the use of new prognostic factors like Ki-67 for the differentiation between luminal A and luminal B or commercial multigene assays like Oncotype DX® (Genomic Health, Redwood City, CA, USA), EndoPredict® (Myriad Genetics, Salt Lake City, UT, USA), Prosigna® (NanoString Technologies, Seattle, WA, USA), or MammaPrint® (Agendia, Irvine, CA, USA), a bigger part of these patients would not be treated with chemotherapy according to current knowledge. If a patient is not a candidate for chemotherapy, according to his/her risk profile, we would not expect to see any difference between the effects of an established standard or an intense dose-dense regimen.

This hypothesis is emphasized by the fact that (intense) dose-dense chemotherapy was effective in hormone receptor-positive patients only in the GIM-2 and AGO-iddETC trials but not in CALGB B9741 (fig. 3). A high-risk situation may effectively erase the (intense) dose-dense and HER2/estrogen receptor (ER) interaction.

Dose-Dense Concepts without G-CSF Support (Weekly Schedule) versus Standard-Dose Chemotherapy

The ECOG 1199/Intergroup trial compared the efficacy of 2 different taxanes, docetaxel and paclitaxel, given either q3w or weekly, in the adjuvant treatment of breast cancer patients. In the 5-year analysis [13], the group receiving 4 cycles of AC followed by 12 weekly cycles of paclitaxel 80 mg/m² showed significantly improved DFS (HR 1.27; p = 0.006) and OS (HR 1.32; p = 0.01) as compared to patients receiving the former standard of paclitaxel 175 mg/m² q3w. Especially patients with HER2-negative disease who received weekly paclitaxel had improved DFS (HR 1.33; p = 0.009) and OS (HR 1.34; p = 0.03), irrespective of their hormone receptor status.
As a consequence of this and other studies [14, 15], weekly paclitaxel became a preferred standard regimen, especially given sequentially after 4 cycles of AC. Doxorubicin 60 mg/m² can be substituted with epirubicin 90 mg/m² [16].

It is important to note that weekly paclitaxel 80 mg/m² is not only dose-dense, but it represents an intense dose-dense schedule. 3-weekly cycles amount to a total dose of 240 mg/m², which leads to a higher total dose and dose intensity in comparison with paclitaxel 175 mg/m² q3w.

Recently, long-term outcome data were reported [17]. When compared with the standard paclitaxel q3w arm, after a median follow-up of 12.1 years, only DFS (HR 0.84; p = 0.01) but no longer OS (HR 0.87; p = 0.09) was statistically different between the weekly paclitaxel arm and the docetaxel q3w arm (HR 0.79; p = 0.001 and HR 0.86; p = 0.054). Clinical outcome and response may vary depending on the breast cancer subtype. For the 2,879 patients with ER-positive/HER2-negative disease, docetaxel q3w was associated with improved DFS (HR 0.76; p = 0.004) but not OS. For the paclitaxel weekly arm, no significant results were observed for either DFS or OS after 12 years of follow-up.

The new analysis also revealed that only paclitaxel weekly was associated with an approximately 30% reduction in the risk of recurrence and death in the triple-negative subtype. Weekly paclitaxel may become a preferred regimen for triple-negative breast cancer.

**Do We Know the Optimal Schedule of Dose-Dense Chemotherapy?**

The SWOG SO221 [18] trial had an open-label 2×2 factorial design to test 2 hypotheses: i) 6 cycles of AC (60/600 mg/m²) q2w versus a novel continuous schedule of intravenous doxorubicin 24 mg/m² once per week, cyclophosphamide 60 mg/m² orally once per day for 15 weeks, supported either by pegfilgrastim or filgrastim; and ii) paclitaxel 80 mg/m² once weekly is superior to 6 cycles of paclitaxel 175 mg/m² q2w.

At a median follow-up of 6 years, an interaction between the 2 randomized factors had emerged so that the 2 randomizations could not be analyzed independently. For the comparison of 6 cycles AC q2w followed by 6 cycles paclitaxel q2w versus 6 cycles AC q2w followed by 12 cycles paclitaxel weekly, the authors reported a trend in DFS (HR 1.24; p = 0.072) and a significant difference in OS (HR 1.46; p = 0.01) favoring patients who received all treatments q2w. Examination of relevant biologic subsets reveals that this advantage for treatment q2w is observed only in patients with hormone receptor-/HER2-negative disease. This observation is the result of a subset analysis that was not protocol-specified and should be regarded as hypothesis-generating.

It must be noted that 6 cycles of paclitaxel q2w were given, rather than the 4 cycles commonly used. This means that no direct comparison can be drawn between 12 weeks of once-weekly paclitaxel and 4 cycles of q2w treatment. Clinically relevant toxic effects, such as allergic-type reactions, musculoskeletal pain, and neuropathy, are more common with the q2w schedule [18].

To what extent the dose-dense application of AC (EC) contributes to the superior efficacy observed in the GIM-2 and CALGB 9741 studies [3] also remains an open question because no study has compared AC (EC) q2w with AC (EC) q3w, followed by weekly paclitaxel in both groups.

A direct comparison between iddETC and 8 cycles of dose-dense ACɉP [3] or AC q3w ×4 followed by weekly paclitaxel ×12 is also missing.

Considering the results of iddETC, one should be aware that dose intensity may also play a role in optimizing the effectiveness of dose-dense regimens.

**Phase III Trials Comparing Different Dose-Dense Concepts**

Three randomized phase III trials have compared different dose-dense regimens. All 3 trials incorporated a 4th cytostatic compound in their experimental arm. The NSABP B-38 trial [9] and the GIM-2 trial [4] intended to improve the dose-dense standard regimen, AC (EC) q2w ×4 followed by paclitaxel q2w ×4 [3]. While NSABP B-38 performed a subrandomization of +/- gemcitabine, the GIM-2 trial evaluated 5-FU as a 4th compound. The GAIN trial intended to improve the results of iddETC by incorporating capecitabine as a 4th cytostatic compound [19]. iddETC was randomized against 4 cycles of EC followed by 10 cycles of paclitaxel weekly in combination with 4 cycles of capecitabine 2,000 mg/m² days 1–14 ×4. The total doses of epirubicin and paclitaxel were identical in both arms, whereas the dosage of cyclophosphamide was lower in the experimental arm.

Hematological and nonhematological toxicity was higher with the 4-drug dose-dense regimens. However, all 3 trials failed to improve the effectiveness of the established (intense) dose-dense 3-drug regimens by incorporating one of these compounds. Hence, 3-drug dose-dense regimens remain the standard of care.

**Conclusion**

Dose-dense chemotherapy should be the standard of care in high-risk breast cancer patients. Dose-dense regimens with 4 drugs have not shown superiority over 3-drug regimens. The importance of total dose in addition to a purely dose-dense concept remains an open question.

**Disclosure Statement**

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