Optimizing the treatment of bevacizumab as first-line therapy for human epidermal growth factor receptor 2 (HER2)-negative advanced breast cancer: an updated meta-analysis of published randomized trials

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Background: Manifold data have demonstrated that the addition of bevacizumab to chemotherapy improved progression-free survival (PFS), while few trials have revealed its significant overall survival (OS) benefit. Furthermore, it still remains suspended how to maximize the benefits of bevacizumab as first-line therapy for human epidermal growth factor receptor 2 (HER2)-negative breast cancer. We sought to conduct a meta-analysis to assess the benefits of bevacizumab with chemotherapy and to identify the ideal chemotherapy partner of bevacizumab in the first-line setting for HER2-negative advanced breast cancer patients.

Methods: Computerized and manual searches were performed to identify randomized clinical trials evaluating the efficacy of bevacizumab plus chemotherapy versus chemotherapy alone or bevacizumab with different chemotherapy regimens as first-line therapy for HER2-negative locally recurrent or metastatic breast cancer patients. Risk ratios or odds ratios with their 95% CIs were used to estimate the association between multiple combinations of bevacizumab with chemotherapy and various clinical outcomes.

Results: With 7 trials identified, this analysis included 3,984 eligible patients. The addition of bevacizumab to chemotherapy resulted in a statistically significant improvement in PFS (P=0.019) and objective response rate (ORR; P<0.001) rather than in OS (P=0.783) when compared with chemotherapy alone. The greater benefits in PFS and ORR were achieved from bevacizumab plus taxane-based regimens compared with bevacizumab plus capecitabine-based regimens, while bevacizumab plus capecitabine had comparable OS with bevacizumab plus paclitaxel. Additionally, bevacizumab-based triplet therapy failed to improve the clinical outcomes when compared with doublet therapy.

Conclusion: This meta-analysis reveals that the addition of bevacizumab to chemotherapy yielded PFS and ORR benefits in HER2-negative advanced breast cancer. Additional studies are still prompted to further optimize the first-line treatment of bevacizumab.

Keywords: breast cancer, bevacizumab, first-line, HER2-negative, meta-analysis

Introduction
Breast cancer is the most common cancer that affects women worldwide. Although its prognosis has been greatly improved due to the rapid development of multidisciplinary therapy, breast cancer still remains the second leading cause of cancer-related...
bevacizumab and chemotherapy in such setting. Furthermore, there are no clear answers on how to maximize the benefits of bevacizumab as first-line therapy for HER2-negative advanced breast cancer. Therefore, we conducted a meta-analysis of all eligible studies to address the overall benefits of bevacizumab as first-line therapy in HER2-negative advanced breast cancer. Additionally, we aimed to determine the best combination of bevacizumab and chemotherapy in such setting.

Methods

Search strategy

The keywords “bevacizumab” or “Avastin”, “phase III” or “phase 3” and “breast cancer” were employed to search the PubMed, Web of Science, EMBASE and EBSCO databases from January 2000 to March 2017 for relevant randomized controlled clinical trials published in English. The selection of the included publications was independently performed by 2 experienced authors (Cunfu Li and Aizhai Xiang). The reference lists of textbooks, previous reviews and all retrieved articles were scrutinized to identify additional studies with potential eligibility. Abstracts, conference information, editorials, letters and case reports were excluded.

Eligibility criteria

We evaluated the full article of each relevant study and included 7 phase III randomized controlled trials that evaluated the efficacy of combining bevacizumab with chemotherapy versus chemotherapy alone or combining bevacizumab with different chemotherapy regimens as first-line therapy for HER2-negative locally recurrent or metastatic breast cancer. Phase I, phase II and nonrandomized studies were excluded. Trials were considered ineligible if they recruited patients to assess neoadjuvant or adjuvant bevacizumab or to evaluate bevacizumab with other target therapies. For multiple reports of the same cohort, the most recent publication was chosen for this analysis.

Data extraction

Data were extracted from eligible publications independently by 2 authors (Xianzhi Chen and Kai Yin). Discrepancies were discussed with a third author (Jinsong Lu), and agreements were reached with consensus according to the Quality of Reporting of Meta-Analyses guidelines.

The following data were extracted from all selected articles: first author, publication year, sample size, number of patients in each arm, bevacizumab and chemotherapy regimens, and either the number of endpoint events in each arm or hazard ratios (HRs) with 95% CIs in each arm.

Statistical analysis

In this meta-analysis, the primary endpoint was PFS, defined as the time from the date of randomization to the date of first documented disease progression or death. The secondary endpoints included OS and objective response rate (ORR), defined as the time from the date of randomization to the date of death from any cause and the percentage of patients who achieved a complete or partial response confirmed ≥28 days after initial documentation of response.

The risk ratio (RR) with its 95% CI was calculated for PFS and OS, while the odds ratio (OR) with 95% CI was calculated for ORR. For the trials with only HRs available, the HRs were used as RRs to facilitate the calculation. The heterogeneity of the study outcomes was calculated using Cochran’s $Q$ statistics ($\chi^2$ test) and $I^2$ test. Either a $P$-value $<0.05$ or $I^2>50\%$ was considered as statistically significant heterogeneity, which indicated the use of a random-effects (DerSimonian and Laird method) model. If there was no between-study heterogeneity, a fixed-effects (Mantel–Haenszel method) model was applied.

Publication bias was evaluated using funnel plots and Begg’s test. Sensitivity analyses were performed to...
quantify the impact of individual trials on the overall effect. A two-sided *P*-value <0.05 was considered significant. All statistical analyses were conducted with Stata statistical software package (release 12.0; Stata Corporation, College Station, TX, USA).

**Results**

**Characteristics of eligible trials**

Considering the quality of the data, we included all the potentially eligible phase III randomized clinical trials with available information for the target population. Based on the search strategy, 7 eligible trials with a total of 3,984 patients were identified and included in this meta-analysis (Figure S1). The details of these included studies are summarized in Table 1. Three of these studies were used to evaluate the efficacy of adding bevacizumab to chemotherapy, including 1,558 women who received bevacizumab combined with chemotherapy and 896 women who were administered chemotherapy alone. The other 4 trials were obtained to assess the optimal chemotherapy partner of bevacizumab. All of them were published as full articles and described as Phase III randomized controlled trials. The former 3 trials were double blinded, and the latter 4 studies were open label. Of note, the E2100 trial was considered ineligible for this meta-analysis despite its inclusion in the

**Table 1** Characteristics of eligible studies

<table>
<thead>
<tr>
<th>Study name</th>
<th>Arms of the study</th>
<th>Primary endpoint</th>
<th>Secondary endpoint</th>
</tr>
</thead>
<tbody>
<tr>
<td>AVADO</td>
<td>Bevacizumab 7.5 mg/kg plus docetaxel 100 mg/m² (day 1); bevacizumab 15 mg/kg plus docetaxel 100 mg/m² (day 1); placebo plus docetaxel 100 mg/m² (day 1); every 3 weeks</td>
<td>Progression-free survival</td>
<td>Best overall response, duration of response, time to treatment failure, overall survival, safety, quality of life</td>
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<tr>
<td>RIBBON-I</td>
<td>Bevacizumab 15 mg/kg (day 1) plus chemotherapy; placebo (day 1) plus chemotherapy; every 3 weeks</td>
<td>Progression-free survival</td>
<td>Overall survival, 1-year survival rate, objective response rate, duration of objective response, progression-free survival based on independent review committee assessment, safety</td>
</tr>
<tr>
<td>MERIDIAN</td>
<td>Bevacizumab 10 mg/kg (days 1, 15) plus paclitaxel 90 mg/m² (days 1, 8, 15); placebo (days 1, 15) plus paclitaxel 90 mg/m² (days 1, 8, 15); every 4 weeks</td>
<td>Investigator-assessed progression-free survival in the intent-to-treat and pVEGF-A&lt;sub&gt;E&lt;sub&gt;populations</td>
<td>VEGF-A-by-treatment interaction test for progression-free survival in the intent-to-treat population, investigator-assessed objective response rate in patients with measurable disease at baseline, duration of objective response in responding patients with measurable disease at baseline, overall survival, 1-year overall survival rate, safety</td>
</tr>
<tr>
<td>TABEA</td>
<td>Bevacizumab 15 mg/kg (day 1) plus taxanes/ capecitabine; bevacizumab 15 mg/kg (day 1) plus taxanes; every 3 weeks</td>
<td>Progression-free survival</td>
<td>Response rate and duration, clinical benefit rate, 3-year overall survival, progression-free survival in patients aged ≥65 years, toxicity, compliance</td>
</tr>
<tr>
<td>SAKK 24/09</td>
<td>Bevacizumab 10 mg/kg every 2 weeks plus paclitaxel 90 mg/m² (days 1, 8, 15) every 4 weeks; bevacizumab 10 mg/kg every 2 weeks plus cyclophosphamide 50 mg/ capecitabine ≥500 mg daily</td>
<td>Incidence of prespecified grade 3–5 adverse events occurring during the trial or within 30 days of the last treatment, regardless of the causal relationship to the trial drug</td>
<td>Objective response rate, disease control rate, progression-free survival, overall survival, other adverse events, quality of life, pharmacoeconomics</td>
</tr>
<tr>
<td>TURANDOT</td>
<td>Bevacizumab 10 mg/kg (day 1, 15) plus paclitaxel 90 mg/m² (days 1, 8, 15) every 4 weeks; bevacizumab 15 mg/kg (day 1) plus capecitabine 1,000 mg/m² twice daily (days 1–14) every 3 weeks</td>
<td>Overall survival</td>
<td>Proportion of patients achieving an overall response, progression-free survival, time to response, duration of response, time to treatment failure, safety, patient-reported outcomes</td>
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<tr>
<td>CARIN</td>
<td>Bevacizumab 15 mg/kg (day 1) plus capecitabine 1,000 mg/m² twice daily (days 1–14); bevacizumab 15 mg/kg (day 1) plus capecitabine 1,000 mg/m² twice daily (days 1–14)/vinorelbine 25 mg/m² (days 1, 8); every 3 weeks</td>
<td>Progression-free survival</td>
<td>Overall response rate, overall survival, safety</td>
</tr>
</tbody>
</table>

**Abbreviations:** VEGF, vascular endothelial growth factor; pVEGF, plasma VEGF.
previous reports on this topic.\textsuperscript{21,22} It recruited both HER2-negative and HER-2 positive patients, but the data were only available for the overall population rather than the HER2-negative subgroup. Therefore, the E2100 trial was excluded in this meta-analysis.

The AVADO trial was a 3-arm trial, in which HER2-negative locally recurrent or metastatic breast cancer patients were randomized on a 1:1:1 basis to receive bevacizumab 7.5 mg/kg plus docetaxel 100 mg/m\textsuperscript{2}, bevacizumab 15 mg/kg plus docetaxel 100 mg/m\textsuperscript{2} or placebo plus docetaxel 100 mg/m\textsuperscript{2} on day 1 every 3 weeks as first-line therapy.\textsuperscript{12} The evaluation of the 2 bevacizumab arms was separately compared with the control arm.

In the RIBBON-1 trial, HER2-negative locally recurrent or metastatic breast cancer patients were randomly assigned in the 2:1 ratio to either bevacizumab 15 mg/kg plus chemotherapy or chemotherapy alone. Chemotherapy was chosen by investigators, including capecitabine (1,000 mg/m\textsuperscript{2} oral twice daily for 14 days), taxane-based (nab-paclitaxel 260 mg/m\textsuperscript{2}, docetaxel 75–100 mg/m\textsuperscript{2}) or anthracycline-based (fluorouracil 500 mg/m\textsuperscript{2}, epirubicin 90–100 mg/m\textsuperscript{2} and cyclophosphamide 500 mg/m\textsuperscript{2}; fluorouracil 500 mg/m\textsuperscript{2}, doxorubicin 50 mg/m\textsuperscript{2} and cyclophosphamide 500 mg/m\textsuperscript{2}; doxorubicin 50–60 mg/m\textsuperscript{2} and cyclophosphamide 500–600 mg/m\textsuperscript{2}; or epirubicin 90–100 mg/m\textsuperscript{2} and cyclophosphamide 500–600 mg/m\textsuperscript{2}) regimen administered every 3 weeks. All enrolled patients were separately analyzed in the capecitabine cohort and the pooled taxane/anthracycline cohort.\textsuperscript{13}

The MERE\textsuperscript{3}DAN trial randomized locally recurrent or metastatic breast cancer patients to paclitaxel 90 mg/m\textsuperscript{2} on days 1, 8 and 15 with or without bevacizumab 10 mg/kg on days 1 and 15 every 4 weeks. The co-primary endpoints were investigator-assessed PFS in the intent-to-treat (ITT) and plasma VEGF-A\textsubscript{high} populations.\textsuperscript{17}

The TABEA study compared taxanes (paclitaxel 80 mg/m\textsuperscript{2} on days 1, 8 and 15 or docetaxel 75 mg/m\textsuperscript{2} on day 1 every 3 weeks) plus bevacizumab (15 mg/kg on day 1 every 3 weeks) with (TBX group) or without capecitabine (1,800 mg/m\textsuperscript{2} daily on days 1–14 every 3 weeks; TB group) as first-line therapy in locally advanced or metastatic breast cancer patients.\textsuperscript{18} In this meta-analysis, TBX and TB were referred to as capecitabine- and taxane-based regimens, respectively.

As to the SAKK 24/09 study, metastatic or locally recurrent inoperable HER2-negative breast cancer patients were randomized to bevacizumab 10 mg/kg every 2 weeks with either paclitaxel 90 mg/m\textsuperscript{2} on days 1, 8 and 15 every 4 weeks or cyclophosphamide 50 mg plus capecitabine 3×500 mg daily. Of note, the primary endpoint was the incidence of prespecified grade 3–5 adverse events (AEs), while PFS, OS and ORR were the secondary endpoints.\textsuperscript{19}

The TURANDOT study was a non-inferiority trial that randomly assigned HER2-negative locally recurrent or metastatic breast cancer patients to either bevacizumab 10 mg/kg on days 1 and 15 plus paclitaxel 90 mg/m\textsuperscript{2} on days 1, 8 and 15 every 4 weeks or bevacizumab 15 mg/kg on day 1 plus bevacitabine 1,000 mg/m\textsuperscript{2} twice daily on days 1–14 every 3 weeks. The primary endpoint was OS, and the secondary endpoints included ORR and PFS. OS was compared in both ITT and per-protocol (PP) populations.\textsuperscript{20} The OS data in the ITT and PP populations were separately included in the corresponding analysis of our study. On the other hand, the primary objective of this trial was to show non-inferior OS with bevacizumab plus capecitabine versus bevacizumab plus paclitaxel in the PP population by rejecting the null hypothesis of inferiority (HR \textgreater;1.33) at a one-sided significant level of 0.025 using a stratified Cox proportional hazard model.\textsuperscript{20} Therefore, the prespecified boundary for HR varied between this trial and the others, which might elicit misleading results. Accordingly, the OS data from this study were also excluded to avoid this confounding effect in this meta-analysis. Herein, there were altogether 3 types of pooled RRs for OS by using different datasets.

The CARIN trial randomized HER2-negative locally recurrent or advanced breast cancer patients to receive bevacitabine 1,000 mg/m\textsuperscript{2} twice daily on days 1–14 plus bevacizumab 15 mg/kg on day 1 every 3 weeks with or without vinorelbine 25 mg/m\textsuperscript{2} on days 1 and 8 every 3 weeks. Despite its superior nature, the significant level was set at 0.05, two sided.\textsuperscript{19}

**Efficacy of adding bevacizumab to chemotherapy**

In terms of the comparison between bevacizumab combined with chemotherapy and chemotherapy alone, there was significant between-study heterogeneity in the RR for PFS (heterogeneity $\chi^2$, 27.15; $I^2$, 85.3%; $P<0.001$; Figure 1). Through the random-effects model, we found a marked improvement in PFS with the addition of bevacizumab 1,000 mg/m\textsuperscript{2} twice daily on days 1–14 plus bevacizumab 15 mg/kg on day 1 every 3 weeks with or without vinorelbine 25 mg/m\textsuperscript{2} on days 1 and 8 every 3 weeks. Despite its superior nature, the significant level was set at 0.05, two sided.\textsuperscript{19}
(OR, 0.560; 95% CI, 0.475–0.661; P<0.001; Figure S3) rather than in OS (RR, 0.986; 95% CI, 0.891–1.090; P=0.783; Figure S2).

Efficacy of bevacizumab plus capecitabine-based chemotherapy compared with bevacizumab plus taxane-based chemotherapy

When it came to the comparison between bevacizumab with capecitabine- and taxane-based chemotherapy, significant between-study heterogeneity failed to be discerned in the RRs for PFS (heterogeneity $\chi^2$, 0.85; $I^2$, 0.0%; $P=0.654$; Figure S4) and OS ([excluding the OS data from the TURANDOT trial: heterogeneity $\chi^2$, 0.05; $I^2$, 0.0%; $P=0.829$; Figure S5]; [including the OS data from the TURANDOT trial in the ITT population: heterogeneity $\chi^2$, 0.17; $I^2$, 0.0%; $P=0.918$; Figure S6]; [including the OS data from the TURANDOT trial in the PP population: heterogeneity $\chi^2$, 0.10; $I^2$, 0.0%; $P=0.951$; Figure S7]) as well as in the OR for ORR (heterogeneity $\chi^2$, 1.31; $I^2$, 0.0%; $P=0.519$; Figure S8). Therefore, the fixed-effects model was used to analyze the data, and a deleterious effect of bevacizumab plus capecitabine-based chemotherapy on PFS (RR, 1.190; 95% CI, 1.103–1.283; $P<0.001$; Figure S4) and ORR (OR, 1.897; 95% CI, 1.535–2.344; $P<0.001$; Figure S8) compared with bevacizumab plus taxane-based regimen was found. Intriguingly, absolute gains in ORR and PFS translated into OS benefit when the data of the TURANDOT trial in the PP population were included (RR, 1.108; 95% CI, 1.005–1.221; $P=0.040$; Figure S7). However, these gains in ORR and PFS were lost no matter whether the data from the TURANDOT trial were excluded (RR, 1.139; 95% CI, 0.880–1.474; $P=0.322$; Figure S5), or its ITT population data were included (RR, 1.091; 95% CI, 0.992–1.201; $P=0.074$; Figure S6).

Efficacy of bevacizumab-based doublet therapy compared with bevacizumab-based triplet therapy

There was significant between-study heterogeneity in both RR for PFS (heterogeneity $\chi^2$, 8.43; $I^2$, 76.3%; $P=0.015$; Figure S9) and OR for ORR (heterogeneity $\chi^2$, 6.80; $I^2$, 70.6%; $P=0.033$; Figure S10) rather than in the RR for OS (heterogeneity $\chi^2$, 2.23; $I^2$, 10.4%; $P=0.327$; Figure 2). Through the random-effects model, bevacizumab-based triplet therapy did not significantly improve the PFS (RR, 1.046; 95% CI, 0.856–1.279; $P=0.658$; Figure S9) or ORR (OR, 0.697; 95% CI, 0.459–0.958; $P=0.090$; Figure S10) when compared with bevacizumab-based doublet therapy. Similarly, no statistical significance was observed for OS with the random-effects model (RR, 0.951; 95% CI, 0.864–1.048; $P=0.311$; Figure 2).

Publication bias and sensitivity analysis

To assess the association between the addition of bevacizumab to chemotherapy and various clinical outcomes in...
HER2-negative locally recurrent or metastatic breast cancer patients, either graphic inspection of funnel plots or quantitative evaluation using Begg’s test indicated the absence of publication bias in OS and ORR but not for PFS ($P$ = 0.014).

As to the studies evaluating the efficacy of bevacizumab plus different types or different numbers of chemotherapy agents, no publication bias was detected with either graphic inspection of funnel plots or quantitative evaluation of Begg’s test in PFS, OS and ORR (data not shown). The sensitivity analyses clarified for each comparison that no individual study affected the overall RRs for PFS and OS as well as the overall ORs for ORR, because omission of any single study made no material difference.

**Discussion**

This meta-analysis is, to the best of our knowledge, the largest and latest study focusing on how to optimize the treatment of bevacizumab as first-line therapy in HER2-negative advanced breast cancer patients.

Our findings indicated that bevacizumab plus chemotherapy significantly improved the PFS and ORR, compared with chemotherapy alone. However, no significant advantage in OS was observed, which was similar to the results in each individual trial. Although the OS is regarded as a gold standard of endpoints and PFS as a surrogate endpoint in metastatic breast cancer, debates still continue for the importance of PFS.\(^{23-25}\) Advocates of OS argue that no therapeutic benefit is achieved without improving OS. This is also the reason why the FDA revoked bevacizumab’s approval for metastatic breast cancer. Using OS instead of PFS as a primary endpoint is relatively difficult and expensive in enrolling the patients and completing the study. Furthermore, it is clinically meaningful to get details on whether certain interventions are capable of delaying disease progression and improving quality of life, which is generally presented by PFS. By this token, advocates of PFS argue that PFS as a primary endpoint has its own merit and plays an important role in the evaluation of a new therapy for metastatic breast cancer, while it is not suitable for a promising agent to attach too much importance on OS.

Notwithstanding the ongoing arguments, researchers have long expected that substantial OS benefit could be obtained from bevacizumab-based regimens by identifying subgroups with specific biomarkers. A meta-analysis, which focused on the first-line bevacizumab therapy, revealed no significant improvement for OS in the triple-negative subgroup.\(^{22}\) On the other hand, the subsequent retrospective biomarker analysis of the AVADO trial revealed that higher plasma concentration of either VEGF-A or VEGFR-2 was associated with more benefit from bevacizumab irrespective of the bevacizumab dose.\(^{26}\) Unfortunately, the MERiDiAN trial, which was the first study to prospectively evaluate plasma VEGF-A, failed to verify the predictive effect of VEGF-A.\(^{27}\)
Furthermore, when the available samples from the AVADO trial were reassessed by using different VEGF assays, the predictive value of plasma VEGF-A disappeared. Collectively, there is no solid evidence to support any biomarker as a predictive factor of bevacizumab efficacy.

This meta-analysis also demonstrated that bevacizumab combined with taxane-based chemotherapy greatly improved the PFS and ORR when compared with bevacizumab plus capecitabine-based regimen. Since the OS analysis of the TURANDOT trial was quite different from that of the other eligible trials, we separately pooled the RRs by excluding the data from the TURANDOT trial and including its data in the PP or ITT population. Only when its data in the PP population were included, did the OS comparison between the 2 groups exhibit statistical significance. Nevertheless, it is necessary for us to be aware of the prespecified boundary for HR in the TURANDOT trial. Given this situation, such advantage has to be interpreted with caution. If we take a close look at the OS data of each included trial, it is obvious that no significant OS benefit was verified in all but one of the included randomized phase III studies. The TURANDOT trial was the first study to compare bevacizumab plus paclitaxel and bevacizumab plus capecitabine. Before this study, these 2 bevacizumab-containing regimens, compared with chemotherapy alone, had both revealed superior PFS and ORR in the previous reports. Unfortunately, the oncologists had no idea whether patient tolerability could be improved by bevacizumab plus capecitabine without compromising the OS in comparison with bevacizumab plus paclitaxel at that time, so they might have been stuck in a dilemma between these two regimens. The TURANDOT trial was conducted to tackle this dilemma. Its final OS analysis met the primary endpoint of the trial, that is, the criterion for non-inferiority in the stratified analysis of the PP population. This finding was also supported by the stratified OS analysis of the ITT population despite the inferiority of bevacizumab plus capecitabine to bevacizumab plus paclitaxel for PFS.

Therefore, the TURANDOT trial provides the evidence that bevacizumab with either paclitaxel or capecitabine has comparable OS in the first-line setting for HER2-negative locally recurrent or metastatic breast cancer patients. Besides, the SAKK 24/09 trial also gives further support for the combined effect of capecitabine with bevacizumab. Although the primary endpoint in this trial was the incidence of pre-specified grade 3–5 AEs instead of ORR, PFS and OS, its results might also offer some clues. The SAKK 24/09 trial showed no statistical significance between bevacizumab plus paclitaxel and bevacizumab plus metronomic capecitabine-cyclophosphamide in terms of ORR, PFS and OS.

Taken together, these findings clarify that the combination of bevacizumab and capecitabine may function as a valid first-line treatment option. In Europe, this regimen, along with bevacizumab plus paclitaxel, has been identified as one of the first-line indications for HER2-negative locally recurrent or metastatic breast cancer patients. However, bevacizumab combined with paclitaxel is the only regimen listed in the latest version of the National Comprehensive Cancer Network Clinical Practice Guidelines in Oncology for Breast Cancer. Furthermore, Delaloge et al demonstrated on the basis of a large-scale real-life setting database that HER2-negative metastatic breast cancer patients who received paclitaxel plus bevacizumab as first-line chemotherapy had a significantly better OS and PFS than those receiving paclitaxel alone. With the accumulation of evidence, we will get a clearer picture of how to choose the preferable chemotherapy partner of bevacizumab.

Last but not the least, when it came to the comparison between the doublet versus the triplet, the addition of a third agent appears to make no sense in the improvement of clinical outcomes. Similar findings have also been reported. The BCIRG 007 trial is a phase III randomized study conducted to compare docetaxel/carboplatin/trastuzumab (TCH) with docetaxel/trastuzumab (TH) as first-line therapy in HER2-amplified metastatic breast cancer. Interestingly, adding carboplatin to TH did not elicit any significant improvement in time to progression, response rate or OS. In many cases, therefore, less is more.

**Conclusion**

This meta-analysis indicated that the combination of bevacizumab and chemotherapy as first-line treatment significantly improved the PFS and ORR in locally recurrent or metastatic breast cancer patients. Greater benefits in PFS and ORR were observed in bevacizumab plus taxane-based regimens compared with bevacizumab plus capecitabine-based ones. In view of the non-inferiority for OS, however, increasing evidence supports the use of bevacizumab plus capecitabine as a preferable first-line option in the USA. Nevertheless, it is far from the end of the story. Additional studies are necessary to further optimize the first-line treatment of bevacizumab.

**Acknowledgment**

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Author contributions

WY and JL conceived and designed the work. CL and AX performed the publication search and the statistical analysis. XC and KY extracted the data. All authors interpreted the results. CL and AX wrote the manuscript draft. All authors participated in the revision. All authors approved the final version.

Disclosure

The authors report no conflicts of interest in this work.

References

Supplementary materials

**Figure S1** Selection process for randomized controlled clinical trials included in the meta-analysis.

**Abbreviation:** RCTs, randomized controlled trials.

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<tr>
<th>Author</th>
<th>Study</th>
<th>Regimen</th>
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<th>RR (95% CI)</th>
<th>% weight</th>
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<td>Mies et al</td>
<td>AVADO</td>
<td>B7.5 + DOC vs PLA + DOC</td>
<td>2010</td>
<td>1.06 (0.88, 1.29)</td>
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<td>B + PAC vs PLA + PAC</td>
<td>2017</td>
<td>0.88 (0.71, 1.09)</td>
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**Figure S2** Forest plot of RR for the association between the addition of bevacizumab to chemotherapy and overall survival in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients.

**Notes:** The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate RR > 1, and symbols on the left of the solid line indicate RR < 1. The combined RR is calculated by the fixed-effects model.

**Abbreviations:** B, bevacizumab; B7.5, bevacizumab 7.5 mg/kg; B15, bevacizumab 15 mg/kg; CAP, capecitabine; chemo, chemotherapy; DOC, docetaxel; PAC, paclitaxel; PLA, placebo; RR, risk ratio; TAX/ANTHRA, taxanes/anthracyclines.
Figure S3 Forest plot of OR for the association between the addition of bevacizumab to chemotherapy and objective response rate in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients.

Notes: The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate OR >1, and symbols on the left of the solid line indicate OR <1. The combined OR is calculated by the fixed-effects model.

Abbreviations: B, bevacizumab; B7.5, bevacizumab 7.5 mg/kg; B15, bevacizumab 15 mg/kg; chemo, chemotherapy; CAP, capecitabine; DOC, docetaxel; OR, odds ratio; PAC, paclitaxel; Pla, placebo; TaX/anThra, taxanes/anthracyclines.

Figure S4 Forest plot of RR for the association between bevacizumab plus different types of chemotherapy agents and progression-free survival in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients.

Notes: The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate RR >1, and symbols on the left of the solid line indicate RR <1. The combined RR is calculated by the fixed-effects model.

Abbreviations: B, bevacizumab; B7.5, bevacizumab 7.5 mg/kg; B15, bevacizumab 15 mg/kg; chemo, chemotherapy; CAP, capecitabine; DOC, docetaxel; OR, odds ratio; PAC, paclitaxel; Pla, placebo; TaX/anThra, taxanes/anthracyclines.
**Figure S5** Forest plot of RR for the association between bevacizumab plus different types of chemotherapy agents and overall survival in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients, excluding the data of the TURANDOT trial.

**Notes:** The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate RR > 1, and symbols on the left of the solid line indicate RR < 1. The combined RR is calculated by the fixed-effects model.

**Abbreviations:** B, bevacizumab; Pac, paclitaxel; rr, risk ratio; TB, taxane/bevacizumab; TBX, taxane/bevacizumab/capecitabine; Xc, capecitabine/cyclophosphamide.

<table>
<thead>
<tr>
<th>Author</th>
<th>Study</th>
<th>Regimen</th>
<th>Publication year</th>
<th>RR (95% CI)</th>
<th>% weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Luck et al(^1)</td>
<td>TABEA</td>
<td>TBX vs TB</td>
<td>2015</td>
<td>1.10 (0.73, 1.65)</td>
<td>39.82</td>
</tr>
<tr>
<td>Rochlitz et al(^3)</td>
<td>SAKK 24/09</td>
<td>B + XC vs B + PAC</td>
<td>2016</td>
<td>1.17 (0.84, 1.63)</td>
<td>80.18</td>
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<tr>
<td>Overall (I^2=0.0%, P=0.829)</td>
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<td></td>
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<td>1.14 (0.88, 1.47)</td>
<td>100</td>
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</tbody>
</table>

**Figure S6** Forest plot of RR for the association between bevacizumab plus different types of chemotherapy agents and overall survival in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients, including the data of the TURANDOT trial in the intent-to-treat population.

**Notes:** The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate RR > 1, and symbols on the left of the solid line indicate RR < 1. The combined RR is calculated by the fixed-effects model.

**Abbreviations:** B, bevacizumab; caP, capecitabine; Pac, paclitaxel; rr, risk ratio; TB, taxane/bevacizumab; TBX, taxane/bevacizumab/capecitabine; Xc, capecitabine/cyclophosphamide.
Figure S7 Forest plot of RR for the association between bevacizumab plus different types of chemotherapy agents and overall survival in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients, including the data of the TURANDOT trial in the per-protocol population.

Notes: The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate RR > 1, and symbols on the left of the solid line indicate RR < 1. The combined RR is calculated by the fixed-effects model.

Abbreviations: B, bevacizumab; caP, capecitabine; Pac, paclitaxel; rr, risk ratio; TB, taxane/bevacizumab; TBX, taxane/bevacizumab/capecitabine; Xc, capecitabine/cyclophosphamide.

Figure S8 Forest plot of OR for the association between bevacizumab plus different types of chemotherapy agents and objective response rate in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients.

Notes: The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate OR > 1, and symbols on the left of the solid line indicate OR < 1. The combined OR is calculated by the fixed-effects model.

Abbreviations: B, bevacizumab; CAP, capecitabine; PAC, paclitaxel; OR, odds ratio; TB, taxane/bevacizumab; TBX, taxane/bevacizumab/capecitabine; Xc, capecitabine/cyclophosphamide.
First-line bevacizumab for HER2-negative advanced breast cancer

<table>
<thead>
<tr>
<th>Author</th>
<th>Study</th>
<th>Regimen</th>
<th>Publication year</th>
<th>RR (95% CI)</th>
<th>% weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Luck et al</td>
<td>TABEA</td>
<td>TBX vs TB</td>
<td>2015</td>
<td>1.23 (0.99, 1.53)</td>
<td>29.32</td>
</tr>
<tr>
<td>Rochlitz et al</td>
<td>SAKK 24/09</td>
<td>B + XC vs B + PAC</td>
<td>2016</td>
<td>1.09 (0.88, 1.34)</td>
<td>29.72</td>
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<td>Welt et al</td>
<td>CARIN</td>
<td>B + CAP/VIN vs B + CAP</td>
<td>2016</td>
<td>0.91 (0.83, 0.99)</td>
<td>40.96</td>
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<tr>
<td>Overall (I²=76.3%, P=0.015)</td>
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<td>1.05 (0.86, 1.28)</td>
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</tbody>
</table>

**Figure S9** Forest plot of RR for the association between bevacizumab plus different numbers of chemotherapy agents and progression-free survival in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients.

**Notes:** The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate RR >1, and symbols on the left of the solid line indicate RR <1. The combined RR is calculated by the random-effects model. Weights are from the random-effects analysis.

**Abbreviations:** B, bevacizumab; CAP, capecitabine; PAC, paclitaxel; RR, risk ratio; TB, taxane/bevacizumab; TBX, taxane/bevacizumab/capecitabine; VIN, vinorelbine; XC, capecitabine/cyclophosphamide.

<table>
<thead>
<tr>
<th>Author</th>
<th>Study</th>
<th>Regimen</th>
<th>Publication year</th>
<th>OR (95% CI)</th>
<th>% weight</th>
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<tr>
<td>Rochlitz et al</td>
<td>SAKK 24/09</td>
<td>B + XC vs B + PAC</td>
<td>2016</td>
<td>1.35 (0.71, 2.60)</td>
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<tr>
<td>Welt et al</td>
<td>CARIN</td>
<td>B + CAP/VIN vs B + CAP</td>
<td>2016</td>
<td>0.63 (0.46, 0.88)</td>
<td>37.76</td>
</tr>
<tr>
<td>Robert et al</td>
<td>RIBBON-1</td>
<td>B + TAX/ANTHR vs B + CAP</td>
<td>2011</td>
<td>0.53 (0.39, 0.70)</td>
<td>39.81</td>
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<tr>
<td>Overall (I²=70.6%, P=0.033)</td>
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<td>0.70 (0.46, 1.06)</td>
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</tbody>
</table>

**Figure S10** Forest plot of OR for the association between bevacizumab plus different numbers of chemotherapy agents and objective response rate in human epidermal growth factor receptor 2-negative locally recurrent or metastatic breast cancer patients.

**Notes:** The size of the square box is proportional to the weight that each study contributes in the meta-analysis. The overall estimate and CI are marked by a diamond. Symbols on the right of the solid line indicate OR >1, and symbols on the left of the solid line indicate OR <1. The combined OR is calculated by the random-effects model. Weights are from the random-effects analysis.

**Abbreviations:** B, bevacizumab; CAP, capecitabine; OR, odds ratio; PAC, paclitaxel; VIN, vinorelbine; TAX/ANTHR, taxanes/anthracyclines; XC, capecitabine/cyclophosphamide.

**References**


