Adjuvant trastuzumab regimen for HER2-positive early-stage breast cancer: a systematic review and meta-analysis

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ABSTRACT

Objective: Breast cancer remains to be the globally leading female cancer. About 15% to 20% of breast cancers have human epidermal growth factor receptor 2 (HER2)-positive tumors – a more aggressive breast cancer subtype with shortened survival. In the light of new and updated trial data on trastuzumab therapy for HER2-positive early-stage breast cancer (EBC), we conducted a systematic review and meta-analysis to update the pooling of its relative treatment effects.

Methods: Systematic search was performed through Pubmed and Scopus to identify studies comparing survival outcomes and risks of heart toxicity effects of adjuvant trastuzumab with chemotherapy versus chemotherapy alone for HER2-positive EBC patients.

Results: Based on the eight included studies in the review, combining trastuzumab with chemotherapy continues to show lowered death and relapse risks by one-third. The decision to initiate trastuzumab, however, needs to be prudently deliberated as two to three times more cardiotoxicity risk was shown to be associated with its use.

Conclusion: Administering adjuvant trastuzumab in a weekly cycle concurrently with anthracycline-taxane chemotherapy regimen appears to be a preferable option to optimize its favorable effect in improving DFS and to prevent significantly higher risk for cardiotoxic effects.

1. Introduction

Breast cancer remains to be the leading cancer among women across the world with approximately 1.67 million cases and 521,907 deaths in 2012 as for the GLOBOCAN cancer incidence, mortality and prevalence report [1]. The World Health Organization reported that in 2008, almost 50% (691,300/1,383,500) of breast cancer cases and 58% (268,900/458,400) of breast cancer deaths occurred in less developed countries, contrary to the notion that breast cancer was a disease of developed countries [2]. They further reported that survival rates for breast cancer varied greatly worldwide, i.e., 80%, 60%, and 40% survival rates in high-income, middle-income and in low-income countries, respectively [3]. Limitations in detection and treatment facilities in less developed countries resulted in a large proportion of women with late diagnosis and thus treatment management; thereby contributing to the poor prognosis in these countries [2].

Human epidermal growth factor receptor 2 (HER2) is a tyrosine kinase receptor that facilitates signaling pathways of cell growth, division, motility and repair [4]. Women with HER2-positive cells are more likely to have a more aggressive form of breast cancer, thus they tend to increase the risk of disease progression and decrease overall survival (OS) [4,5]. Hence, the determination of HER2 positivity through immunohistochemical (IHC) assay or in situ hybridization (ISH) assay became clinically important. The prevalence of HER2-positive breast cancer in Western population was approximately 15% to 20% of primary breast cancers [6], which was not much different compared to the estimated HER2 positivity rate in Asian population which was about 23.5% [7].

Currently, the 2018 American Society of Clinical Oncology (ASCO) [8], the 2017 US National Comprehensive Cancer Network (NCCN) Guideline on Breast Cancer [9], and the 2015 European Society for Medical Oncology (ESMO) Clinical Practice Guidelines for the Diagnosis, Treatment, and Follow-Up of Primary Breast Cancer [10] recommended the administration of trastuzumab (the first monoclonal antibody-based therapy developed to specifically target HER2) with chemotherapy for the management of HER2-positive early-stage breast cancer (EBC) in adjuvant settings. Trastuzumab may be administered by 3-week cycle (i.e., initial dose of 8 mg/kg, followed by 6 mg/kg every 3 weeks) or by weekly cycle (i.e., Initial dose 4 mg/kg, scheduled weekly.).
Trastuzumab has been recommended for the treatment of HER2-positive breast cancer but its associated with cardiotoxic effects. An updated systematic review and meta-analysis along with exploration on the impact of prognostic factors and intervention characteristics, and net clinical benefit was performed in aid of establishing a recommended optimum treatment regimen for adjuvant trastuzumab for HER2-positive patients. Current evidence available to date indicated that combining adjuvant trastuzumab with chemotherapy might gain benefits. Current evidences available to date indicated that combining adjuvant trastuzumab with chemotherapy might significantly improve the OS and DFS of HER2-positive EBC women thereby decreasing the risk for mortality and relapse by one-third, but also increases the risk for cardiotoxic effects by two to three times more. Combining adjuvant trastuzumab with chemotherapy might gain benefits of DFS and OS about one-fifth and one-third or more than CHF. Administering adjuvant trastuzumab in a weekly cycle concurrently with anthracycline-taxane chemotherapy regimen appears to be a preferable option to optimize its favorable effect in improving DFS and to prevent significantly higher risk for cardiotoxic effects.

Followed by 51 further weekly doses of 2 mg/kg, and sequentially or concurrently with standard chemotherapy of anthracycline-taxane regimens (e.g., doxorubicin and cyclophosphamide (AC) plus paclitaxel/docetaxel; or, fluorouracil, epirubicin and cyclophosphamide (FEC) plus docetaxel/paclitaxel), taxane regimens (e.g., docetaxel plus cyclophosphamide), or anthracyline regimen (e.g., AC). The treatment is recommended to start within two to six weeks after the surgery [9,10].

Trastuzumab is currently available as 150 mg (single-dose vial) and 440 mg (multi-dose vial) powder for concentrate solution for intravenous (IV) infusion; and for subcutaneous (SC) injection containing 600 mg/5mL. It works by binding to the juxtamembrane domain of HER2 thereby triggering the downregulation of HER2 expression. However, the treatment effect of adjuvant trastuzumab among EBC women with HER2-positive in improving the patient’s disease-free survival (DFS) and OS was controversial, as significant adverse cardiac effects (i.e., congestive heart failure, CHF and left ventricular ejection fraction, LVEF decline) were reported [11–13].

As for our knowledge, there were seven meta-analyses (N = 4–6 randomized controlled trials, RCTs) [14–20] and three systematic reviews only (N = 5–6 RCTs) [21–23]. Results of the seven meta-analyses [14–20] indicated that adjuvant trastuzumab with chemotherapy in treating HER2-positive EBC patients significantly improved the OS and DFS when compared to chemotherapy alone with a reduction in the risk of mortality ranging from about 30% to 34%, and a reduction in the risk of recurrence ranging from 34% to 50%. However, none of them comprehensively explored which prognostic factors (i.e., cycle regimen, type of administration, and the combined chemotherapy type) might affect both the relative treatment efficacy and safety. Results of this may aid in establishing a recommended optimum treatment regimen for adjuvant trastuzumab for HER2-positive EBC patients. In addition, one new RCT and two updated RCT results have been published since the last meta-analysis. We therefore conducted a systematic review and meta-analysis to update the pooling of the relative treatment efficacy and safety of adjuvant trastuzumab plus chemotherapy compared to chemotherapy alone in HER2-positive EBC patients. Subgroup analysis by prognostic factors and intervention characteristics was explored.

2. Methods

2.1. Search strategy

The review protocol was developed following the Preferred Reporting Items for Systematic reviews and Meta-Analyses (PRISMA) guidelines [24] and was registered at PROSPERO with registration number CRD42017072583. Electronic searching for relevant studies published from inception to 8 July 2017 was performed through two electronic databases (i.e. Medline via PubMed and Scopus), without language restrictions using search terms and strategies described in Table S1 (Supporting Information).

2.2. Selection of studies

Two review authors (AJG and DTO) independently screened all the titles and abstracts that were identified in the systematic searching. The full text of potentially eligible studies with relevant abstracts and titles were retrieved and independently evaluated for eligibility using the set inclusion and exclusion criteria. All excluded studies were recorded and properly accounted for reasons for exclusion. Any disagreement between the selection results of review authors was resolved by discussion and consensus with the supervisor (UC).

2.3. Inclusion and exclusion criteria

RCTs and non-RCTs were included if they met the following criteria: participating patients were HER2-positive EBC covering stage I, IIA, IIB, and IIIA; assessed the relative treatment effects of adjuvant trastuzumab plus any chemotherapy regimen (i.e., anthracycline-taxane regimen, anthracycline-only regimen, taxane-only regimen) regardless of the cycle regimen type, timing and duration of administration; compared with any chemotherapy alone regimen; and, assessed any of following primary outcomes, i.e., overall survival (OS) defined as the time since randomization to death from any cause, disease-free survival (DFS) defined as the time since randomization to the date of recurrence of tumor; and/or secondary outcomes, i.e., congestive heart failure (CHF) defined as New York Heart Association as class III-IV, and left ventricular ejection fraction (LVEF) decline which was defined as absolute decline of at least 10% from baseline LVEF or a decline to less than 50% or 55%.

Studies were excluded if they evaluated: among populations which included/combined local and metastatic HER2-positive breast cancer patients; trastuzumab in neoadjuvant trastuzumab settings; trastuzumab with other anti-HER2 drugs; trastuzumab compared with different dosages or duration of the same interventions; and, if data are insufficient for pooling after contacting the authors.
2.4. Data extraction and management

We constructed and used data extraction forms which were composed of five data domains including study characteristics, patient characteristics (e.g., demographic data and relevant cancer data), interventions (e.g., dosing and cycle regimen type, duration of therapy, type of trastuzumab administration with chemotherapy), chemotherapy comparator, and outcomes data (e.g., type of outcomes, frequency data, and summary statistics). Data under these key domains were extracted and recorded from each included study. Any inconsistency in the extraction of data or in the assessment of studies was resolved by consensus with the supervisor (UC).

2.5. Risk of bias assessment

Included RCTs were assessed for Risk of Bias using Cochrane Handbook for Systematic Reviews of Interventions [25], composed of the following domains: random sequence generation, allocation sequence, blinding, incomplete outcome data and selective outcome reporting. Each domain was rated as low risk if relevant methods and procedures being assessed based on the criteria are adequate; high risk if procedures are inadequate; and unclear risk if unclear or not explicitly reported in the manuscript. Further, observational studies were assessed for quality using the Newcastle Ottawa Scale for Quality Assessment [26], which is composed of the following domains: selection of cohorts, comparability of cohorts and outcome assessment. Each domain includes item questions complemented with a scoring algorithm to rate each domain as good, fair or poor depending on the score that the study attained by meeting the set criteria for low risk of bias. The studies were then rated as having a low, moderate or high risk of bias if the percentage score was ≥75%, 50–74% and <50%, respectively. Two reviewers (AJG and DTO) independently assessed each study using the tools and pre-specified criteria. The supervisor (UC) was consulted for any disagreement in the ratings of the assessment.

2.6. Data synthesis

HR was used to assess relative treatment effects on OS and DFS, while RR was used to measure the safety outcomes CHF and LVEF decline. A meta-analysis of aggregated data from published studies was performed to pool the relative treatment effects using a random-effects model by DerSimonian and Laird method if heterogeneity was present (i.e., Cochran’s Q test <0.10 or $I^2 > 25\%$); otherwise, a fixed-effects model by inverse variance method was used. Sources of heterogeneity were further explored by fitting the co-variables (i.e., tumor size, nodal, hormone-receptor status, type of intervention, trastuzumab duration, regimen cycle, administration type, length of follow-up period and the risk of bias) one-by-one in a meta-regression model. A decrease in $I^2$ by at least 50% would suggest that such co-variable might be a source of heterogeneity. Sub-group analysis was then performed if data were available and sufficient.

A net clinical benefit was applied to assess risk (i.e., CHF and LVEF decline) and benefit (i.e., OS and DFS) simultaneously [27]. For this, risk difference (i.e., incremental ($\Delta$)) was estimated for each study and pooled across studies using a random-effects model. An incremental risk and benefit ratio (IRBR) was then estimated. In addition, data for $\Delta$risk and $\Delta$benefit were further simulated using Monte Carlo with 1,000 replications assuming normal distributions for both $\Delta$risk and $\Delta$benefit. Finally, a cost-effective plane curve was constructed assigning $\Delta$risk on Y-axis and $\Delta$benefit on X-axis.

All the analyses were performed using STATA software version 14 and Microsoft excel version 2016. Two-sided $P < 0.05$ was considered statistically significant except for the subgroup analysis and heterogeneity test, in which $P < 0.10$ was used. Funnel plot and Egger’s test were used to detect potential publication bias. If any of these suggested asymmetry, a contour-enhanced funnel plot was constructed to further explore the likely attributable source of such asymmetry.

3. Results

3.1. Search results

Figure 1 illustrates the study flow of this systematic review. The combined electronic search from Medline and Scopus provided a total of 2,878 records. After removing the duplicates, 2,672 studies were remained for the screening of titles and abstracts. Of these, 2,602 studies were excluded leaving 65 studies plus one additional study from reference lists, for full text assessment. Of these, 58 studies did not meet the inclusion criteria, leaving eight studies included in the review [28–35].

3.2. Study characteristics

Study and patient characteristics of included studies are summarized in Table 1. All the included studies were RCTs. None of the detected non-RCTs met our criteria because their patients in the comparator group were mixed with chemotherapy and no treatment. Of these, three studies [32–34] were conducted in Europe, two studies [28,35] were international multicentre trials with collaborated countries from the different regions, and three studies [29–31] were from USA. The median follow-up period ranged from 3.9 to 11 years. The number of participating HER2-positive EBC women ranged from 232 to 4,046 with median age of 48 to 62 years. Majority of the included patients had a tumor size of more than 2 cm, while only three studies [28,32,33] provided information on the proportion of patients whose tumor size is less than 1 cm but with no report on their corresponding outcomes. The percentage of node-positive patients ranged from 52% to 100%, and hormone-receptor (HR)-positive patients were 40% to 73%. The percentage of patients who underwent mastectomy ranged from 36% to 67%.

Information on intervention and comparator characteristics are summarized in Table S2 and S3 (Supporting Information). Four studies [29–33] administered trastuzumab in a 3-week cycle while three studies [28,34,35] implemented the weekly cycle. Courses of treatment were 12 months for seven studies [28–32,34,35] and 9 weeks for one study [33]. Trastuzumab was administered with chemotherapy concurrently in five studies [29–33] and sequentially in three studies [28,34,35]. Note that concurrent use of trastuzumab in our review refers to its concomitant use with...
3.3. Risk of bias assessment of included studies

Risk of bias assessment was performed and results were described in Figure S4 (Supporting Information). Among eight studies, most studies, 5/8 [28–31,33] were rated as low risk for random sequence generation (N = 5 [28–31,33]), selective reporting (N = 7 [28–32,34,35]), and incomplete outcomes (N = 5 [28–31,33]). However, most studies (N = 6 [28–32,34]) were unclear whether they had applied allocation concealment, whereas none of the studies applied blinding.

3.4. Overall Survival (OS)

The OS was obtained from six RCTs [28,29,32–35] which included a total of 5,355 and 5,280 of HER2-positive EBC women in the intervention and the control groups, respectively. Among these two groups, 736 and 1,023 patients died with pooled death rates and 95% confidence interval (CI) of 11.9% (8.5%, 15.5%) and 16.7% (11.4%, 22.0%). The reported
The DFS was estimated from six studies [28,29,32–35], analyzing a total of 10,635 HER2-positive EBC women. The total number of recurrences was, respectively, 1,271 out of 5,355 and 1,676 out of 5,280 in the trastuzumab-chemotherapy and chemotherapy alone groups with pooled recurrence rates (95% CI) of 21.6% (16.6%, 26.5%) and 29.4% (24.6%, 34.2%). The relative treatment effects of trastuzumab-chemotherapy regimen versus chemotherapy alone regimen on recurrence were moderately heterogeneous with the $I^2$ of 61.1% as shown in Figure 3. The pooled HR was 0.65 (95% CI: 0.55, 0.75, $P < 0.001$), suggesting significantly lower risk of recurrence in the trastuzumab-chemotherapy group, about 35% when compared to the chemotherapy alone group. Possible sources of heterogeneity in the pooling of HRs were then explored by performing subgroup analysis and among the pre-identified covariates, trastuzumab cycles (i.e., 3-week versus weekly cycles), intervention type (i.e., chemotherapy regimen combined with trastuzumab) and nodal status were found to be associated with the reduction in the degree of heterogeneity in some subgroups. For the subgroup analysis based on trastuzumab cycles, three RCTs [28,34,35] applied 3-week cycle and three RCTs [29,32,33] administered by weekly cycle. The results for the subgroup analysis based on cycle regimen apply for the subgroup analysis based on administration type as the trials belonging to the subgroups of the 3-week cycle and weekly cycle are similar for the subgroup status were found to be associated with the reduction in the degree of heterogeneity in some subgroups. For the subgroup analysis based on trastuzumab cycles, three RCTs [28,34,35] applied 3-week cycle and three RCTs [29,32,33] administered by weekly cycle. The results for the subgroup analysis based on cycle regimen apply for the subgroup analysis based on administration type as the trials belonging to the subgroups of the 3-week cycle and weekly cycle are similar for the subgroups of sequential and concurrent administration, respectively. The pooling within the subgroups of 3-week cycle versus weekly cycle yielded the $I^2$ values of 73.9% and 0%, respectively, as shown in Figure S5 (Supporting Information). However, the pooled HRs for these corresponding groups were not much different with the pooled HRs of 0.68 (95% CI 0.44–0.91, $P < 0.001$) for the 3-week cycle and 0.61 (95% CI 0.55–0.68 $P < 0.001$) for the weekly cycle.

For the subgroup analysis based on intervention type, four RCTs [29,32,33,35] administered trastuzumab with anthracycline-taxane chemotherapy regimens while two RCTs [28,34] administered trastuzumab with any/mixed chemotherapy regimens (i.e., anthracycline-taxane chemotherapy, anthracycline-based chemotherapy, or taxane-based chemotherapy). The pooling of subgroups of trials which used trastuzumab with anthracycline-taxane chemotherapy regimen versus trastuzumab with any/mixed chemotherapy regimen type resulted in $I^2$ values of 0% and 0%, respectively, as shown in Figure S6 (Supporting Information). The pooled subgroup HR for trastuzumab with any/mixed chemotherapy regimen type was significantly higher (HR 0.77; 95% CI: 0.68, 0.85, $P < 0.001$) compared to the pooled
subgroup HR for trastuzumab with anthracycline-taxane chemotherapy regimen (HR 0.60; 95% CI: 0.54, 0.66, P < 0.001). This indicates that higher reduction in the risk of recurrence was observed in the intervention arm of trials which have administered trastuzumab with anthracycline-taxane chemotherapy regimens (i.e., 40%), as compared to the reduction in the risk of recurrence from the intervention arm of the trials which have administered trastuzumab with any/mixed chemotherapy regimen type (i.e., 23%).

For the subgroup analysis based on nodal status, five RCTs [29,32–35] involved 60% or more node-positive patients in their study while only one trial by Cameron et al., 2017 [28] involved less than 60% node-positive patients. The pooling of subgroup of trials involving 60% or more node-positive patients decreased the $I^2$ to 25.5%, as shown in Figure S7 (Supporting Information). The pooled subgroup HR for trials which involved 60% or more node-positive patients was lower (HR 0.61; 95% CI: 0.53, 0.70, P < 0.001) compared to the HR of one trial with less than 60% node-positive patients (HR 0.76; 95% CI: 0.67, 0.85, P < 0.001) suggesting higher reduction in the risk of recurrence observed in the administration of trastuzumab among trials where more node-positive patients have participated (i.e., 39%), as compared to the reduction...
in the risk of recurrence of the trial with less participating node-positive patients (i.e., 24%).

The funnel plots for OS and DFS are provided in S8 and S9 (Supporting Information). The funnel plots and Egger’s tests for OS and DFS showed symmetry, both with non-statistically significant p values for Egger’s test, suggesting no evidence of publication bias.

3.6. Congestive Heart Failure (CHF)

The data on CHF were obtained from six RCTs [28,30–34] analyzing a total of 4,779 and 4,879 HER2-positive EBC women in the trastuzumab-chemotherapy and the chemotherapy alone groups, respectively. A total of 100 and 79 patients of these corresponding groups developed CHF with pooled rates (95% CI) across studies of 2.0% (1.1%, 2.9%) and 0.6% (0.2%, 0.9%). The reported risk ratios (RRs) and 95% CIs of individual studies and showed generally higher CHF events in the intervention group compared to the control group except for Joensuu et al., 2009 [33], with a degree of heterogeneity of 0%. The pooled RR for CHF was 3.71 (95% CI: 2.41, 5.71, P < 0.001) as shown in Figure 4, which could be interpreted that the risk of CHF in the trastuzumab-chemotherapy group increased by 3.71 times more compared to the chemotherapy alone group.

3.7. Left Ventricular Ejection Fraction (LVEF) decline

Data for LVEF decline were extracted from five trials [28,32–35] analyzing a total 3,327 and 3,262 HER2-positive EBC women in the trastuzumab-chemotherapy and the chemotherapy alone groups, respectively. Among these two groups, 308 and 148 developed LVEF decline with pooled rates across studies of 8.4% (2.5%, 14.2%) and 4.7% (0.7%, 8.8%). The reported RRs and 95% CIs of individual studies similarly showed that LVEF decline events were generally higher in the intervention group compared to the control group except for Joensuu et al., 2009 [33]. The relative treatment effects of trastuzumab-chemotherapy regimen versus chemotherapy alone regimen on the risk of LVEF decline were substantially heterogeneous with the I² of 80.8%. The pooled RR was 2.17 (95% CI: 1.11, 4.24, P < 0.001), as presented in Figure 5, suggesting a significantly higher risk of LVEF decline in the trastuzumab-chemotherapy group by 2.17 times more when compared to chemotherapy alone group.

Possible sources of heterogeneity in the pooling of RR for LVEF decline were then explored and among the pre-identified covariates, trastuzumab cycle regimen, administration type, and intervention type (i.e., chemotherapy regimen combined with trastuzumab) were found to be associated. For the subgroup analysis of LVEF decline based on trastuzumab cycle regimen, three RCTs [28,34,35] administered by 3-week cycle and two RCTs [32,33] using the weekly cycle. Similarly, the results for the subgroup analysis based on cycle regimen apply for the subgroup analysis based on administration type. The pooling within subgroups of 3-week cycle versus weekly cycle resulted in the I² values of 15.5% and 69.2%, respectively, as presented in Figure S10 (Supporting Information). The pooled RR for the 3-week cycle was significantly higher (RR 4.16; 95% CI: 2.50, 6.92, P < 0.001) compared to the pooled RR for the weekly cycle (RR 1.24; 95% CI: 0.54, 2.83, P = 0.614).

For the subgroup analysis of LVEF decline based on intervention type, three RCTs [28,32–35] administered trastuzumab with anthracycline-taxane chemotherapy regimen while two RCTs [28,34] administered trastuzumab with any/mixed chemotherapy regimen type yielding the I² values of 40% and 0%, respectively, as shown in Figure S11 (Supporting Information). The pooled RRs of these two corresponding subgroups were 4.70 (95% CI: 2.98, 7.41, P < 0.001) and 1.32 (95% CI 0.74–2.36, P = 0.350), respectively.

3.8. Risk and benefit assessment

Net clinical benefits were assessed to simultaneously compare risks (i.e., CHF and LVEF decline) and benefits (i.e., OS and DFS).

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**Figure 4. Forest plot for overall RR of CHF.**
Incremental risks and benefits were estimated for individual included studies and then pooled across studies using a random-effects model, see Figure S12 (Supporting Information). IRBRs were further estimated, i.e., 0.1895, 0.3005, 0.5135, and 0.8144 for ΔCHF/ΔDFS, ΔCHF/ΔOS, ΔLVEF decline/ΔDFS, and ΔLVEF decline/ΔDFS, respectively (see Table S13 [Supporting Information]); the lower IRBR reflects the better benefit over the risk. Comparing with CHF, the IRBRs were about 1/5 and 1/3, i.e., DFS and OS would be one-fifth and one-third or more than the development of CHF. For LVEF decline, the IRBRs were a half and four-fifth, i.e., DFS and OS were about 50% and 20% or lower than the development of LVEF decline. Monte Carlo with 1000 replications was applied to simulate data Δrisks and Δbenefits, cost-effective plane curves were constructed, see Figure S14 a-d (Supplemental Information). If the threshold is set at 0.5, receiving trastuzumab-chemotherapy should gain benefits from DFS and OS than the risk of CHF, but not for LVEF decline.

4. Discussion

We conducted a systematic review and meta-analysis of eight RCTs which demonstrated that adding adjuvant trastuzumab to chemotherapy compared to chemotherapy alone in HER2-positive EBC patients can reduce the risk of mortality and recurrence of 33% and 35%, respectively. However, adjuvant trastuzumab might result in three and two times increased risk of CHF and LVEF decline compared to using the standard chemotherapy regimens alone. Net clinical benefit was assessed considering risk and benefit simultaneously which showed that DFS and OS were about one-fifth and one-third or more than the development of CHF, whereas DFS and OS were about a half and one-fifth or lower than the development of LVEF decline.

We updated the previous meta-analyses by including new and latest RCTs with the longest follow up period report to date [28–31,35]. Our results were consistent with the results of previous meta-analyses [14–20] which showed risk reduction for death and recurrence by about 30% to 34% and 36% to 50%, respectively. Likewise, our pooled results showed increasing risk of CHF and LVEF decline by about 3.71 and 2.17 times, respectively, which corresponded with previous meta-analyses results of 2.45 to 7.60 times for CHF and 1.88 to 2.45 times for LVEF decline. We, however, have added more evidence that receiving adjuvant trastuzumab therapy might gain benefits of DFS and OS about one-fifth and one-third or more over adverse drug reactions of CHF compared with receiving chemotherapy alone. While our study excluded neoadjuvant settings and focused on trastuzumab adjuvant use as it is the first and more widely used indication especially among developing countries, many patients in developed countries are currently treated with neoadjuvant trastuzumab therapy and dual blockade trastuzumab with pertuzumab.

Hence, judicious decision on the trade-offs between these beneficial and unfavorable treatment effects remain to be an important consideration prior to deciding the initiation of the therapy. Given this result and the fact that trials included in this analysis have all excluded patients with pre-existing cardiovascular disease or low LVEF level (i.e., less than 50%), it is mandatory to assess cardiovascular functions prior to initiation and continuously monitoring during the trastuzumab-chemotherapy therapy.

Moreover, this analysis explored on several factors which may possibly affect both the treatment efficacy and safety outcomes. Among the assessment of factors related to the characteristics of the patients, study and the intervention regimens, the results have shown the association of treatment effect (i.e., DFS and LVEF decline) with the type of intervention regimen particularly the chemotherapy regimen type combined with trastuzumab, the cycle regimen type of trastuzumab, and the type of trastuzumab administration.
The subgroup analyses showed improved DFS or significantly higher reduction in the risk of recurrence when adjuvant trastuzumab was administered in a weekly cycle concurrently with anthracycline-taxane chemotherapy regimens. As regards the type of trastuzumab administration, the results of this updated analysis was consistent with the previous results by Yin et al., 2011 [18] showing better DFS improvement for concurrent administration (OR 0.65, 95% CI 0.49–0.85) compared to sequential administration (OR 0.74, 95% CI 0.65–0.83).

In terms of cardiotoxic outcomes, the subgroup analyses also showed that the risk of LVEF decline appeared to be significantly higher when adjuvant trastuzumab was administered in a 3-week cycle sequentially with any/mixed type of chemotherapy regimens but the risks were dramatically declined with the weekly cycle regimen, concurrent administration and trastuzumab combined with anthracycline-taxane chemotherapy. However, these latter poolings might be limited by the small number of included studies and thus lack of statistical power.

In this light, cardiotoxic outcomes associated with trastuzumab use remains to be an important decision point by clinicians and patients, regardless of the cycle regimen, administration type and intervention regimen. On the basis of these comparative subgroup results weighing both the efficacy and safety outcomes, adjuvant trastuzumab administered in a weekly cycle concurrently with anthracycline-taxane chemotherapy regimens appears to be a preferable option in terms of optimizing its beneficial treatment effect as shown by its higher risk reduction for relapse, and in terms of averting higher risk for detrimental outcomes as shown by its lesser magnitude of risk of LVEF decline.

Further research is recommended to explore and determine which particular chemotherapy regimen type combined with trastuzumab provides the best improvement in survival outcomes and least risk of cardiotoxic effects towards identifying the best treatment regimen for this therapy. Assessment of optimal treatment regimen based on other relevant patient-related outcomes (e.g., quality of life) as well as cost-effectiveness and the role of biosimilars which can lower the cost of therapy to improve its accessibility especially among resource-constrained settings are likewise strongly recommended. Moreover, it is speculative that in the future the related research in this field will evolve to provide clear evidence on the role of adjuvant trastuzumab among HER2-positive patients with small node-negative tumors which could lead to the improvement of clinical practice for these patients.

5. Conclusion

In the light of the most current and longest trial evidence available to date, combining adjuvant trastuzumab with chemotherapy is able to gain the benefits of OS and DFS over the risk of CHF but not for LVEF decline when compared to chemotherapy alone in HER2-positive EBC women. The currently available evidence under the subgroup analyses showed that administering adjuvant trastuzumab in a weekly cycle concurrently with anthracycline-taxane chemotherapy regimen is able to lower cardiogenetic toxicity than 3-week cycle given not much differences in the benefit of OS and DFS.

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Author contribution statement

UC and AT supervised the overall direction of the research. AJG, UC, AT, and TR designed and developed the protocol. AJG and OTD independently performed the systematic search, study selection, data extraction and risk of bias assessment with the supervision of UC and AT. AJG, OTD, and AT performed the statistical analysis. AJG, UC, AT, and TR analyzed and interpreted the results. AJG took the lead in the drafting of the manuscript and all authors contributed in its review and finalization.

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Declaration of interest

The authors have no relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript. This includes employment, consultancies, honoraria, stock ownership or options, expert testimony, grants or patents received or pending, or royalties.

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