Immune checkpoint inhibitor-related adverse cardiac events in patients with lung cancer: a systematic review and meta-analysis

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Abstract

Background: Although people are more and more aware of the cardiotoxicity caused by immune checkpoint inhibitors (ICIs) in the treatment of lung cancer, its incidence rate has not been systematically analyzed. This study aims to evaluate the incidence of cardiotoxicity related to the ICI therapies for lung cancer, so as to enhance clinicians’ attention to cardiotoxicity, implement proper prevention and intervention for high-risk patients, and minimize the risk of cardiac dysfunction during and after completion of therapy.

Methods: We conducted a systematic literature search for relevant publications in PubMed and Scopus from inception to 19 April 2022. Pooled incidence and risk ratios with 95% confidence intervals (95% CIs) for cardiotoxicity events were calculated.

Results: A total of 37 studies covering 38 trials, including 14,342 patients, were identified. The pooled risk ratios of incidence of any cardiac AEs were 1.944 [95% CI 0.8–4.725] (Single ICI versus chemotherapy), 1.677 [95% CI 1.065–2.64] (Single ICI plus chemotherapy versus chemotherapy), and 0.478 [95% CI 0.127–1.798] (Single ICI versus Dual ICI). The incidence of myocarditis and arrhythmia were 0.003 [95% CI 0.002–0.006] and 0.014 [95% CI 0–0.037], respectively.

Conclusion: Single ICI did not increase the risk of cardiotoxicity compared with chemotherapy, and single ICI plus chemotherapy increased the risk of cardiotoxicity by 67% compared with chemotherapy alone. Combination immunotherapy did not increase the risk of cardiotoxicity compared with single ICI.

Keywords: Immune checkpoint inhibitor, Immunotherapy, Cardiotoxicity, Myocarditis, Immune related adverse event

Introduction

Patients with lung cancer, especially with advanced or metastatic lung cancer, are often poorly treated due to high morbidity and mortality [1]. The treatment prospects of this refractory disease, however, have changed with the in-depth research on immune checkpoint inhibitors (ICIs) in recent years [2]. Immune checkpoints are immunosuppressive molecules that protect human tissues and organs by regulating the immune response to maintain tolerance. They are monoclonal antibodies that prevent these molecules from releasing the immune system and killing tumor cells [3], including PD-1, PD-L1 and CTLA-4. As ICIs are widely used in the treatment of lung cancer, especially metastatic and advanced lung cancer [4], an excessively enhanced immune response has led to a wide range of immune related adverse events, including cardiotoxicity [5] that may be serious and have a poor prognosis, such as myocarditis, pericardial disease [5], non-inflammatory left ventricular dysfunction [6] and

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myocardial infarction (MI) [7]. Adverse cardiac events caused by ICIs occur at a low rate but can be accompanied by life-threatening events. Studies have shown that the mortality of affected patients remains as high as 50% [8, 9]. Although people are more and more aware of the cardiotoxicity caused by ICIs in the treatment of lung cancer, its incidence rate has not been systematically analyzed.

For cancer survivors, asymptomatic or symptomatic treatment related cardiac dysfunction or cardiac abnormalities may be responsible for interruption or discontinuation of cancer-directed therapies, which may reduce the chance for long-term survival [10]. By analyzing all published randomized clinical trials (RCTs) on ICIs, this study aims to evaluate the incidence of cardiotoxicity related to the ICI therapies for lung cancer, so as to enhance clinicians’ attention to cardiotoxicity, implement proper prevention and intervention for high-risk patients, and minimize the risk of cardiac dysfunction during and after completion of therapy.

Methods
The study was registered with INPLASY202250042 (https://inplasy.com/inplasy-2022-5-0042/) and reported in accordance with the PRISMA statement [11].

Search strategy and selection criteria
We conducted a systematic literature search for relevant publications in PubMed and Scopus from inception to 19 April 2022. Review articles, case series, conference abstracts, and articles not published in English were excluded. The full search strategies are supplied in Additional file 1: M1. Additional articles were identified through reference lists and relevant systematic reviews. We considered all randomized studies on ICIs for lung cancer. Studies were eligible if they reported outcome data with regards to immune related adverse events. Observational studies were not considered.

Study selection and data extraction
The study selection and data extraction were performed by two authors independently. Disagreements were resolved through discussion. Data were extracted, including first author, publication year, study design, study registration, treatments, sample size in each arm, tumor type and stage, follow-up time, outcome measures. The primary outcome of this meta-analysis was the risk ratio of any cardiotoxicity between two ICI-related therapies (including Single-ICI vs Chemotherapy, Single-ICI+Chemotherapy vs Chemotherapy, and Single-ICI vs Dual-ICI). The secondary outcomes were incidence of ICI-associated myocarditis, pericardial effusion, heart failure, cardiopulmonary events, cardiac arrest, atrial fibrillation, arrhythmia, and MI. Risks of bias were assessed independently using the Risk of Bias Tool developed by the Cochrane Collaboration [12].

Statistical analysis
The incidence of cardiotoxicity may be very rare, even no event occurring in either or both arms of a study. Meta-analysis of incidence using inverse variance methods has the problem that the variance becomes very small when the incidence is small or large, with the consequence that such studies get a large weight in the meta-analysis. Transformation methods can be used to avoid undue large weight for studies with small or large incidence. The double arcsine transformation [13] has properties that make it the clearly preferred option over the often-used logit transformation. Pooled incidence and risk ratios (RRs) with 95% confidence intervals (95% CIs) for cardiotoxicity events were calculated. This meta-analysis was conducted in MetaXL 5.3 (EpiGear International) using the IVhet (inverse variance heterogeneity) model [14]. The Chi² test and the Higgins I² statistics were used to assess heterogeneity between the included studies [15]. In addition, sensitivity analyses were performed by a leave-one-out analysis. Publication bias was assessed with the LFK index and Doi plot. The Interpretation of the index in terms of asymmetry was in Additional file 1: M2.

Results
Study characteristics
Our literature search returned 1081 articles, of which 315 were assessed as eligible. A total of 37 studies covering 38 trials, including 14,342 patients, were identified to be based on qualitative analyses (Fig. 1). Among them, six trials were phase 1a/b study, fourteen trials were phase 2 study, and eighteen trials were phase 3 study. Nine trials covered patients with small-cell lung cancer (SCLC) and 29 trials reported patients with nonsmall-cell lung cancer (NSCLC). Nineteen trials reported cardiac adverse events (AEs) with single ICI, twelve trials reported cardiac AEs with single ICI plus chemotherapy, and seven trials reported cardiac AEs with dual ICI plus or minus radiotherapy. Seven trials provided data on cardiac AEs of only ICI versus chemotherapy, nine trials provided data on cardiac AEs of single ICI versus single ICI plus chemotherapy, and four trials provided data on single ICI versus dual ICI. The characteristics of each study are shown in Table 1. Additional file 1: Fig. S1 and S2 describe the risk of bias according to each study and a summary of the risk of bias, respectively. Except for five trials, namely KEYNOTE-598 [16], KEYNOTE-189 [17], PACIFIC [18], IMpower133 [19], and CA184-156 [20] was a double-blind trial, the other 32 trials were open...
label trials. The risk of attrition bias exists in seven trials due to small sample size.

**Primary outcomes**

*Single ICI versus chemotherapy*

The pooled RR of incidence of any cardiac AEs across the seven studies was 1.944 [95% CI 0.8–4.725], suggesting that the incidence of any cardiac AEs with single ICI treatment was 1.944 times higher than with chemotherapy, but was statistically insignificant (p-value = 0.142). $I^2$ was 16%, indicating very small heterogeneity. Table 2, Fig. S3, Additional file 1: Table S1.

*Single ICI plus chemotherapy versus chemotherapy*

The pooled RR of incidence of any cardiac AEs across the nine studies was 1.677 [95% CI 1.065–2.64], suggesting that the incidence of any cardiac AEs with single ICI plus chemotherapy was 1.677 times higher than with chemotherapy, which was statistically significant (p-value = 0.026). $I^2$ was 0%, indicating no heterogeneity. Table 2, Additional file 1: Fig. S4, Table S2.

*Single ICI versus dual ICI*

The pooled RR of incidence of any cardiac AEs across the four studies was 0.478 [95% CI 0.127–1.798], suggesting
| First author | Year | Study | Study design | Phase | Tumor type | Treatments | Sample size | Median follow-up (month) |
|--------------|------|-------|-------------|-------|------------|------------|-------------|--------------------------|
| Altorki [21] | 2021 | NCT02904954 | single-centre, open-label, randomised, controlled, | 2 | clinical stages I-IIIA NSCLC | neoadjuvant durvalumab alone versus neoadjuvant durvalumab + stereotactic radiotherapy | 60 | 16.9 |
| Antonia [22] | 2016 | NCT01928394 | multicentre, open-label | 1/2 | limited-stage or extensive-stage SCLC, had disease progression after at least one previous platinum-containing regimen | Nivolumab versus Nivolumab + ipilimumab | 213 | 198.5 days |
| Bang [23] | 2020 | JVDF | single-arm, non-randomised, multi-cohort | la/b | advanced NSCLC | Ramucirumab | 28 | 22.6 |
| Boyer [16] | 2021 | KEYNOTE-598 | randomized, double-blind, | 3 | Metastatic NSCLC PDL1 tumor proportion score $\geq$ 50% | Pembrolizumab versus Pembrolizumab + ipilimumab | 568 | 24.0 |
| Felip [24] | 2020 | - | open-label, multicenter, dose-escalation and expansion | 1b | stage IIIb/IV ALK-rearranged NSCLC | Ceritinib + Nivolumab | 36 | 24.6 |
| Garassino [25] | 2018 | NCT02087423 | open-label, single-arm | 2 | advanced NSCLC | Durvalumab | 444 | 12.0 |
| Gettinger [26] | 2021 | Lung-MAP S1400I | open-label randomized | 3 | previously treated patients with Stage IV squamous Cell Lung Cancer | Nivolumab + ipilimumab versus Nivolumab | 246 | 29.5 |
| Herbst [27] | 2021 | JVDF | multi-cohort, non-randomized, open-label, | 1a/b | treatment-naive, locally advanced unresectable or metastatic NSCLC | ramucirumab + pembrolizumab | 26 | 23.5 |
| Hui [28] | 2017 | KEYNOTE-001 | international, randomized, open-label | 1 | advanced NSCLC | Pembrolizumab | 101 | 22.2 |
| Ikeda [29] | 2020 | TORG1936/AMBITIOUS jRCTs031190084 | multicenter, single-arm | 2 | NSCLC with idiopathic interstitial pneumonias | Atezolizumab | 17 | 3.0 |
| Jotte [30] | 2020 | IMpower131 | global, open-labe | 3 | stage IV squamous NSCLC | Atezolizumab + carboplatin + paclitaxel versus Atezolizumab + carboplatin + nab-paclitaxel | 1000 | 18.1 |
| Juergens [31] | 2020 | NCT02537418 | multcenter multi-cohort | 1b | advanced, metastatic, recurrent or unresectable SCLC | Durvalumab + tremelimumab + chemotherapy | 22 | 19.6 |
| Kanda [32] | 2016 | (JapicCTI)-132,071 | single-center, open-label | 1b | stage IIIb without indication for definitive radiotherapy, stage IV, or recurrent NSCLC | Nivolumab + chemotherapy | 24 | 6 |
| First author | Year | Study | Study design | Phase | Tumor type | Treatments | Sample size | Median follow-up (month) |
|--------------|------|-------|--------------|-------|------------|------------|-------------|------------------------|
| Langer [33]  | 2016 | KEYNOTE-021 NCT02039674 | randomised, open-label | 2 | advanced NSCLC | Pembrolizumab + chemotherapy versus Chemotherapy | 123 | 10.6 |
| Lin [34]     | 2020 | DETTERED NCT02525757 | randomised, open-label | 2 | non-metastatic and unresectable NSCLC | atezolizumab | 40 | 15.3 |
| Malhotra [35]| 2021 | NCT03026166 | multicenter, open-label | 1–2 | Previously Treated Extensive-Stage SCLC | Rova-T + nivolumab versus Rova-T + nivolumab and ilimumab | 42 | 7.3 |
| Mark [36]    | 2021 | SAKK 19/17 | multicenter, single-arm, and open-label trial | 2 | locally advanced, stage IIIB to IV, cytology or histology proven NSCLC | durvalumab | 21 | 6.0 |
| Mazieres [37]| 2021 | POPLAR NCT01903993 | randomized, open-label | 2 | previously treated advanced NSCLC | Atezolizumab versus Docetaxel | 277 | 48.0 |
| Mazieres [37]| 2021 | OAK NCT02008227 | randomized, open-label | 3 | previously treated advanced NSCLC | Atezolizumab versus Docetaxel | 1187 | 48.0 |
| Mok [38]     | 2019 | KEYNOTE-042 NCT02220894 | multicenter, randomized, open-label | 3 | previously untreated, PD-L1-expressing, locally advanced or metastatic NSCLC | Pembrolizumab versus Chemotherapy | 1274 | 12.8 |
| Nishio [39]  | 2021 | IMpower132 NCT02657434 | multicenter, randomized, open-label | 3 | advanced NSCLC | atezolizumab + Chemotherapy versus Chemotherapy | 101 | 17.5 |
| Pakkala [40] | 2020 | NCT02701400 | randomized, two-arm, non-comparative | 2 | relapsed SCLC | durvalumab(D) + tremelimumab(T) without SBRT versus SBRT followed D/T | 18 | 5.7 |
| Ramalingam [41]| 2022 | JASPER | multicenter, open-label | 2 | advanced ( unresectable) or metastatic NSCLC (stage 3B/4) | Niraparib + Pembrolizumab | 38 | 6.0 |
| Rodríguez-Abreu [17]| 2021 | KEYNOTE-189 NCT02578680 | double-blind trial | 3 | metastatic nonsquamous NSCLC without sensitizing EGFR/ ALK alterations | Pembrolizumab + chemotherapy versus Placebo + chemotherapy | 607 | 31.0 |
| Schoenfeld [42]| 2022 | NCT02888743 | open-label, multicentre, randomised | 2 | metastatic NSCLC refractory to previous PD(L)-1 therapy | Durvalumab–tremelimumab ± radi-therapy | 78 | 12.4 |
| Sezer [43]   | 2021 | EMPOWER-Lung 1 NCT03088540 | multicentre, open-label, global | 3 | advanced NSCLC | Cemiplimab versus Chemotherapy | 697 | 13.1 |
| Welsh [44]   | 2020 | NCT02444741 | prospective randomized | 1/2 | metastatic NSCLC | Pembrolizumab with or without radiation therapy | 100 | 20.4 |
| Welsh [45]   | 2020 | – | single-center, open-label | 1/2 | Limited-Stage SCLC | Pembrolizumab and chemoradiation | 40 | 23.1 |
| Antonia [18]| 2017 | PACIFIC | randomized, double-blind, international | 3 | stage III, locally advanced | Durvalumab versus placebo | 709 | 14.5 |
| First author | Year | Study | Study design | Phase | Tumor type | Treatments | Sample size | Median follow-up (month) |
|--------------|------|-------|--------------|-------|------------|------------|-------------|-------------------------|
| Barlesi [46] | 2018 | JAVELIN Lung 200 NCT02395172 | open-label, multicentre, randomised | 3     | stage IIIB, IV, or recurrent NSCLC with disease progression after previous platinum doublet treatment | Avelumab group versus Docetaxel | 758 | 18.9 |
| Borghaei [47] | 2015 | CheckMate-057 NCT01673867 | randomized, open-label, international | 3     | stage IIIB or IV or recurrent nonsquamous NSCLC after radiation therapy or surgical resection | Nivolumab versus Docetaxel | 555 | – |
| Herbst [48]  | 2016 | KEYNOTE-010 | open-label, multicentre, randomised | 2/3   | previously treated, PD-L1-positive, advanced NSCLC | Pembrolizumab versus Pembrolizumab versus Docetaxel | 991 | 13.1 |
| Horn [19]    | 2018 | IMpower133 | double-blind, placebo-controlled | 3     | Extensive-Stage SCLC | Atezolizumab versus Placebo | 394 | 13.9 |
| Paz-Ares [49] | 2019 | CASPIAN NCT03043872 | open-label, multicentre, randomised | 3     | extensive-stage SCLC | Durvalumab + platinum– etoposide versus Platinum– etoposide | 531 | 14.2 |
| Reck [20]    | 2016 | CA184-156 NCT01450761 | multicentre, randomized, double-blind | 3     | Extensive-Stage SCLC | Chemotherapy/Ipilimumab versus Chemotherapy/Placebo | 954 | 10.5 |
| Socinski [50] | 2018 | Impower-150 | international, randomised, open-label | 3     | Metastatic Nonsquamous NSCLC who had not previously received chemotherapy | Atezolizumab + BCP versus Bevacizumab + Carbo + Paclitaxel (BCP) | 787 | 15.5 |
| West [51]    | 2019 | Impower-130 | multicentre, randomised, open-label | 3     | metastatic NSCLC | Atezolizumab + chemotherapy versus Chemotherapy | 705 | 19.2 |
| Carbone [52] | 2017 | CheckMate-026 | multicentre, randomised, open-label | 3     | Stage IV or Recurrent NSCLC | Nivolumab versus Chemotherapy | 530 | 13.5 |
|              |      |       |              |       |             |             | Total 38 studies | 14,342 |

NSCLC nonsmall-cell lung cancer, SCLC small-cell lung cancer
that the incidence of any cardiac AEs with single ICI was 47.8% of that with dual ICI, but was statistically insignificant (p-value = 0.275). I² was 0%, indicating no heterogeneity. Table 2, Additional file 1: Fig. S5, Table S3.

### Secondary outcomes

The incidences of any cardiac AEs with single ICI, single ICI plus chemotherapy, and dual ICI plus or minus radiotherapy were 0.007 [95% CI 0.001–0.015], 0.019 [95% CI 0–0.048], and 0.024 [95% CI 0–0.068], respectively, showing that they were in an increasing trend. Table 2, Additional file 1: Fig. S6–S8. During ICI treatment, the incidence of myocarditis and arrhythmia was 0.003 [95% CI 0.002–0.006] and 0.014 [95% CI 0–0.037], respectively. The incidence of other cardiac damage was shown in Table 2, Additional file 1: Fig. S9–S16.

### Subgroup analyses

We divided the ICI related cardiac AEs into SCLC and NSCLC subgroups for meta-analysis. The incidence of ICI related cardiac AEs in SCLC subgroup was 0.010, while that in NSCLS subgroup was 0.013. Due to the lack of studies comparing ICI related cardiac AEs of these two types of lung cancer, it cannot be explained which subgroup has a significantly higher incidence. Table 2, Additional file 1: Fig. S17.

The subgroup analysis of Single-ICI + chemotherapy vs chemotherapy showed that the RR of cardiac toxicity of ICI + CPA vs CPA was 1.877 [1.121–3.143], indicating that the incidence of cardiotoxicity of ICI + CPA was 1.88 times higher than that of CPA, with statistical difference, while there was no statistical difference between ICI + CPE and CPE, or between ICI + CE and CE. Table 2, Additional file 1: Fig. S4.
**Sensitivity analyses**

We performed a sensitivity analysis of all pooled results using leave-one-out analysis. When PACIFIC [18] was excluded, $I^2$ decreased from 80 to 20%, and the incidence of any cardiac AEs with single ICI treatment decreased from 0.007 to 0.004, indicating that the heterogeneity of the pooled effect size (ES) mainly came from PACIFIC. When KEYNOTE-010 was excluded, $I^2$ dropped from 80 to 0%, the pooled RR of incidence of any cardiac AEs with single ICI versus chemotherapy went from statistically insignificant 1.944[95%CI 0.8–4.725] to statistically significant 2.374 [95%CI 1.158–4.867]. This suggests that the heterogeneity of the pooled ES mainly came from KEYNOTE-010 [48], which altered the statistical significance of the pooled ES. When Impower-130 [51] was excluded, the pooled RR of incidence of any cardiac AEs with single ICI plus chemotherapy versus chemotherapy went from statistically significant 1.677 [95%CI 1.065–2.64] to statistically insignificant 1.257[95%CI 0.585–2.699], suggesting that the pooled ES were sensitive to Impower-130 [51], which altered the statistical significance of the pooled values. No sensitive studies were found in any of the other pooled ES.

**Publication bias**

LFK index showed that there was major asymmetry and significant publication bias for the all results of "pooled incidence" except the pooled incidence of "cardiopulmonary". But the results of three comparisons (single ICI versus chemotherapy, single ICI plus chemotherapy versus chemotherapy, and single ICI versus dual ICI) showed minor asymmetry and publication bias. Additional file 1: Fig. S18–S32. Interpretation of the LFK index in terms of asymmetry see Additional file 1: M2.

**Discussion**

A total of 38 studies involving 14,342 lung cancer patients were included in this meta-analysis. Our findings showed that there was no significant difference in the incidence of cardiotoxicity between single ICIs and chemotherapy alone, and that the increased risk of cardiotoxicity with combination immunotherapy versus single ICIs was statistically insignificant were fully consistent with the meta-analysis performed by Agostineto et al. [53]. This should be good news for lung cancer patients. However, the incidence of cardiotoxicity with single ICIs and combination immunotherapy was 0.7% and 2.4%, respectively. This was also confirmed in an analysis of Vigibase (The World Health Organization's international database of case safety reports) by Salem et al. [9], who observed that among 30,000 cancer patients treated with ICIs, combination immunotherapy exhibited a significantly higher rate of myocarditis (1.33%) than monotherapy did (0.31%). In addition, the mortality of myocarditis secondary to combination immunotherapy was higher than that of monotherapy (67% vs. 36%), suggesting that combination immunotherapy had a more severe myocarditis [54]. Similar findings were reported by Johnson et al. [55] in a query of the Bristol Myers Squibb Company safety database. We think this is due to the large differences of population and intervention between retrospective studies and RCTs. RCT's population is ideal for random assignment into groups that enjoy a similar baseline, whereas a retrospective study population comes from the real world and is susceptible to selection bias, and even if matching is performed, the results can be affected by various biases. “Pure” treatment in the RCT intervention and control groups are guaranteed in the best possible way to avoid exposure to other drugs, but in retrospective study all interventions are performed in clinical settings with a variety of comorbidities. Whatever, it’s a real side of the real world. The results should therefore be interpreted with caution. Simultaneously, our study also found that ICIs plus chemotherapy increased the risk of cardiotoxicity by 67% compared with chemotherapy alone, suggesting that sometimes the combination is more cardiotoxic than monotherapy. Sensitivity analysis suggested that when removing IMpower-130 [51] changes the statistical significance. As can be seen from Figure S4, IMpower-130 is the study with the smallest confidence interval and significant weight in this comparison. So, this study had the largest impact on the pooled effect size. We believe that the weight assigned to IMpower-130 is reasonable using the IVhet model, and we prefer the pooled effect size with IMpower-130.

According to our pooled analysis of various cardiotoxicity, myocarditis showed the lowest incidence (0.3%), cardiac arrhythmia exhibited the highest incidence (1.4%), and the incidence of MI and pericardial effusion was 0.6% and 1.1%, respectively. Although the consequences of myocarditis and MI are serious, the high incidence of arrhythmia and pericardial disease cannot be ignored in clinical setting. Although the ICIs related cardiotoxicity mechanisms are currently unknown, there is a strong association between immune responses and heart disease. Severe systolic dysfunction/heart failure and fatal arrhythmias are often triggered by viral and autoimmune myocarditis. The heart is particularly vulnerable to immune related damage, and immune responses that normally lead to tissue damage and inflammation are particularly dangerous for the heart. The reason for this lies in its dense blood vessels that provide access to antibodies and immune cells, its anatomy is nonredundant and even small lesions can provide a substrate for arrhythmias [56]. Previous studies have demonstrated that PD-L1, PD-1, and CTLA-4 are important signaling
pathways in cardiac immune crosstalk, and abrogation of those pathways leads to autoimmune myocarditis and heart failure [57, 58]. The independent autoantibody is the mechanism by which T cell-mediated responses to cardiac antigens promote disease progression and heart failure through myocardial inflammatory cell infiltration and increased myocardial fibrosis [59]. Collectively, acute MI, ventricular arrhythmias, autoimmune T cell-mediated myocarditis and conduction disease may be triggered by suppressing PD-L1, PD-1, or CTLA-4, and direct inhibition of PD-L1 may inevitably accelerate pre-existing heart disease, and invite noninflammatory cardiomyocyte dysfunction in diseased hearts even in the absence of an immune response.

In view of the severe cardiotoxicity with ICIs, detection of cardiac biomarkers in serum might be useful for baseline-based risk stratification, early diagnosis of cardiovascular disease during and after treatment, and identification of cancer patients who might benefit from cardioprotective therapy during continuing oncological treatment, as well as identifying patients with cardiovascular disease who might require long-term follow-up. Cardiac troponins (cTn) T and cTnI are structural proteins unique to the heart and are therefore organ-specific markers. Troponin assessment can help identify patients who may benefit from preventive treatment for cardiotoxicity and monitor response to cardioprotective therapy. We sought to perform a meta-analysis of serum biomarkers in patients receiving cardiotoxic cancer ICI therapies, by collecting as few as three relevant observational studies that provided some noteworthy results. According to Mahmood et al. [8], among 35 ICI associated myocarditis patients and 105 ICI non-myocarditis patients, those who experienced major adverse cardiac events (MACE) obtained a higher admission, peak, and discharge/final troponin T value than those who did not. Patients with final/ discharge troponin T greater than or equal to 1.5 ng/ml were bound to a fourfold increased risk of MACE. Petricciuolo et al. [60] studied 30 patients who had high-sensitivity troponin T measured before starting ICI therapy. After 3 months of treatment, The MACE occurred only in 7 patients (23%) with high-sensitivity troponin T ≥ 14 ng/L at baseline. However, according to Yuan et al. [61], no significant changes in cTnI were found in a cohort of 19 cancer patients whose biomarkers were assessed at baseline, 1, 3, and 6 months after ICI administration. In our opinion, more studies are needed to determine whether cTn has the potential to be a serum biomarker for cardiotoxicity in ICI patients. Heart failure is a well-recognized complication that impacts survival and quality of life. It’s a progressive disorder [62]. This process begins with cardiotoxicity of immunotherapy and/or chemotherapy, and is usually progresses after structural change of the heart. It is increasingly important to address chronic and long-term adverse treatment effects in cancer survivors. For those high-risk lung cancer survivors, routine monitoring through cardiac imaging may be required after completion of lung cancer-directed therapy, in order to initiate appropriate interventions to prevent or even reverse the progression of cardiac dysfunction [10].

This meta-analysis has several limitations. For ethical reason in cancer treatment, the studies we included were basically from open label trials. In addition, there are also several options for chemotherapy or radiotherapy combined with ICI, and only one study was placebo-controlled. All these factors may lead to inter-study heterogeneity. Due to the extremely low incidence of cardiotoxicity, it is possible that no events will occur in any treatment arm, those studies with no cardiac events in all arms were excluded from pooling, the results of this meta-analysis might be overestimated. Due to the lack of adequate studies, there was no meta-analysis of serum biomarkers of cardiotoxicity in this study and no relevant conclusions were drawn.

**Conclusion**

In summary, our study showed that single ICI did not increase the risk of cardiotoxicity compared with chemotherapy, and single ICI plus chemotherapy increased the risk of cardiotoxicity by 67% compared with chemotherapy alone. Our findings also suggested that combination immunotherapy did not increase the risk of cardiotoxicity compared with single ICI, and the conclusions of this meta-analysis should be interpreted with caution because of inconsistencies with the results of large retrospective studies.

**Abbreviations**

ICIs: Immune checkpoint inhibitors; RCTs: Randomized clinical trials; MI: Myocardial infarction; RRs: Risk ratios; SCLC: Small-cell lung cancer; NSCLC: Nonsmall-cell lung cancer; AES: Adverse events; ES: Effect size; cTn: Cardiac troponins; MACE: Major adverse cardiac events.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s12935-022-02760-2.

**Additional file 1: M1. Search terms on PubMed. M2. Interpretation of the index in terms of asymmetry. Figure S1. Risk of bias summary: review authors’ judgements about each risk of bias item for each included study. Figure S2. Risk of bias graph: review authors’ judgements about each risk of bias item presented as percentages across all included studies. Figure S3. Forest plot of rate ratio of any cardiac adverse events among patients with lung cancer. Single immune checkpoint inhibitor vs Chemotherapy. Figure S4. Forest plot of rate ratio of any cardiac adverse events among patients with lung cancer. Single immune checkpoint inhibitor +Chemotherapy vs Chemotherapy. Figure S5. Forest plot of rate ratio of any cardiac adverse events among patients with lung cancer.
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Author contributions
XTZ and TL drafted the manuscript. NG contributed to the development of the selection criteria, and the risk of bias assessment strategy. ZJX provided statistical expertise. All authors read, provided feedback and approved the final manuscript.

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Availability of data and materials
All data generated or analysed during this study are included in this published article and its Additional information files.

Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

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