Clinicopathological and prognostic significance of PD-L1 expression in sarcoma
A systematic review and meta-analysis

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Abstract
Programmed cell death-ligands 1 (PD-L1) is a key immune checkpoint protein and a promising therapeutic target for malignancy tumor immunotherapy. The prognostic value of PD-L1 in patients with bone and soft tissue sarcoma remains controversial. Therefore, this meta-analysis is conducted to evaluate the associations of PD-L1 expression with overall survival (OS), progression-free survival (PFS), and clinicopathological characteristics of sarcoma.

A comprehensive literature search of PubMed, Web of Science, Embase, and Cochrane Library was conducted for relevant studies. A total of 14 studies published from 2013 to 2017 were included. The pooled hazard ratios (HRs) with 95% confidence intervals (CIs) were extracted from included studies to assess the association between PD-L1 expression and OS, PFS of patients with sarcoma. Other relevant data were extracted to evaluate the correlations of PD-L1 expression with risk and clinicopathological characteristics of sarcoma. Stata 12.0 software was applied to calculate the strength of association between PD-L1 expression and sarcoma.

In total, 14 articles containing 15 independent studies and 1,451 patients were included in this meta-analysis. We found that the high PD-L1 expression was associated with poorer overall survival (HR 1.27, 95% CI: 0.70–1.84 \( P = .000 \)) and poorer events-free survival (HR 2.05, 95% CI: 1.55–2.70, \( P = .000 \)) in bone and soft-tissue sarcoma patients. Additionally, we conducted subgroup analysis according to histology type, ethnicity, target of PD-L1 assessment, cutoff, the significance of correlations with poor overall survival and events-free survival were also observed. In contrast none of the clinicopathological characteristics (gender, age, tumor site, tumor grade, tumor depth, tumor necrosis rate, metastasis, recurrence, chemotherapy, radiotherapy) was found to be associated with PD-L1 expression in our analysis.

The findings from this meta-analysis indicate that PD-L1 expression might be a useful predictive factor of poor prognosis for patients with bone and soft tissue sarcoma.

Abbreviations: CAS = cutaneous angiosarcoma, CI = confidence interval, CS = chondrosarcoma, DFS = disease-free survival, EFS = event-free survival, HR = hazard ratio, MFS = metastasis-free survival, OR = odds ratio, OS = osteosarcoma, OS = overall survival, PFS = progression-free survival, RFS = recurrence-free survival, STS = soft tissue sarcoma.

Keywords: bone and soft tissue sarcoma, meta-analysis, prognosis, programmed cell death ligand 1 (PD-L1)

1. Introduction

Sarcomas are a group of malignant tumors of mesenchymal origin and characterized by heterogeneous subtypes with various cytogenetic profiles. Skubitz and D’Adamo divided sarcomas into 2 general categories: primary bone sarcoma and soft-tissue sarcoma in 2007.\textsuperscript{[1]} In recent years, with the development of surgical techniques and the emergence of effective chemotherapy, combining chemotherapy, radiotherapy with surgery became the standard treatment for sarcoma. However, the overall survival still was stagnant and the 5-years probability of local recurrence, metastasis remained high.\textsuperscript{[2,3]} Meanwhile, the prognosis is poorer in the majority of these cases, the average survival period after developing recurrence and metastasis is less than 1 year.\textsuperscript{[4]} The limitations in current treatment of sarcoma desperately need breakthroughs, particularly for case with metastatic and relapse disease, to treat these devastating diseases.

Immunotherapy is regarded as an emerging therapeutic option in oncology and immune escape is the important biological process for malignant tumor progression.\textsuperscript{[5,6]} Recent studies have proved programmed death 1 (PD-1) and programmed death-ligand 1 (PD-L1) play a pivotal role in the immune surveillance by
2. Results

2.1. Search result and characteristics of studies

In this meta-analysis, we identified a total of 511 potentially relevant articles after the initial search of PubMed, Embase, Cochrane Library, and Web of Science. After screening the titles and abstracts, 472 studies were excluded because of the duplicate or irrelevant. Thirty-nine potentially eligible articles were further reviewed in detail, 25 articles were excluded, including 14 articles without sufficient data, 2 recruiting overlapping patients, 4 articles including patients with other tumors, 5 articles that were not relevant. As a result, 14 articles containing 15 independent studies and 1451 patients were selected for this meta-analysis. The detailed diagram of filtering process is presented in Fig. 1.

The basic characteristics of the eligible studies are summarized in Table 1.[13–26] The sample sizes of each study ranged from 13 to 470 patients. And a total of 1451 patients from Asian, Europe, and America were enrolled in the studies. Because the study was carried out by Koirala, 2 cohorts of patients were enrolled, and PD-L1 was reported independently, so they were analyzed as 2 individual studies. All 15 of the eligible studies for the analysis were retrospectively designed. The proportion of PD-L1 positivity in the above studies ranged from 5.87% to 64.7%. The histological category of tumors included osteosarcoma (6 studies), soft tissue sarcoma (8 studies), and chondrosarcoma (1 study). Five studies were reported on Asians, and 10 studies on Caucasians. The PD-L1 protein expression was detected by immunohistochemistry (IHC) in 12 studies, while the remaining study used quantitative real-time PCR (1 study) and RNA sequencing (2 studies) to survey the PD-L1 mRNA expression. The survival endpoints were reported by different effect size including overall survival (OS), disease-free survival (DFS), metastasis-free survival (MFS), recurrence-free survival (RFS), and events-free survival (EFS). The HR estimations for 6 studies were directly reported, whereas the other studies provided the clinicopathological parameters of sarcoma patients.

Table 1

| Study | Years | Country | Design | Number of patients | Mean age (range) | Median follow-up (range) | Histology type | PDL-1 assessment Target Assay | PDL-1 positivity cutoff | NOS score |
|-------|-------|---------|--------|-------------------|-----------------|-------------------------|---------------|-----------------------------|------------------------|-----------|
| Que et al.[13] | 2017 | China | RC | 163 | 39 (5–77) | 75 (1–178) | STS | Protein | IHC | >1% of tumor cells | 11.70% | 8 |
| Budczies et al.[14] | 2017 | Germany | RC | 162 | NR | NR | STS | mRNA | RNA seq | NR | 21.10% | 8 |
| Sundara et al.[15] | 2017 | Netherlands | RC | 22 | 18 (7–70) | 56 (14–117) | OS | Protein | IHC | >1% of tumor cells | 18.20% | 7 |
| Shen et al.[16] | 2014 | USA | RC | 37 | 29 (6–75) | 36 (1–200) | OS | mRNA | qPCR | NR | 37.50% | 8 |
| Liu et al.[17] | 2017 | China | RC | 72 | NR | NR | OS | Protein | IHC | Total score ≥7 | 41.0% | 6 |
| Kostine et al.[18] | 2015 | Netherlands | RC | 21 | NR | 54 (15–100) | OS | Protein | IHC | >1% of tumor cells | 52.00% | 7 |
| Koirala cohort1[19] | 2016 | USA | RC | 41 | 16 | 54 (15–100) | OS | Protein | IHC | >1% of tumor cells | 29.30% | 8 |
| Kim J et al.[20] | 2013 | Korea | RC | 105 | NR | 35 (1–175) | STS | Protein | IHC | Total score ≥7 | 64.70% | 8 |
| Kim C et al.[21] | 2016 | South Korea | RC | 82 | 26 (1–79) | 33.8 | STS | Protein | IHC | Total score ≥7 | 43.0% | 7 |
| Honda et al.[22] | 2013 | Japan | RC | 106 | 74.5 | 20 (2–100) | CAS | Protein | IHC | >5% of tumor cells | 30.20% | 9 |
| Costa et al.[23] | 2017 | Brazil | RC | 13 | 32.9 (23–65) | 31.8 (2–156) | Oral OS | Protein | IHC | Total score ≥7 | 61.50% | 8 |
| Chowdhury et al.[24] | 2017 | UK | RC | 59 | NR | 33 (3–200) | STS | Protein | IHC | >5% of tumor cells | 59.30% | 8 |
| Bertucci et al.[25] | 2017 | France | RC | 470 | 62 (10–92.12) | NR | STS | mRNA | RNA seq | NR | 41.0% | 6 |
| D’Angelo et al.[26] | 2015 | USA | RC | 47 | 46 (22–76) | NR | STS | Protein | IHC | >1% of tumor cells | 8.50% | 8 |

CAS = cutaneous angiosarcoma, CS = chondrosarcoma, IHC = immunohistochemistry, NOS = Newcastle–Ottawa scale, NR = not reported, OS = osteosarcoma, qPCR = quantitative real-time polymerase chain reaction, RC = retrospective cohort, RNA seq = RNA sequencing, STS = soft tissue sarcoma.

*Total score was calculated by adding a score of staining percentage to another score of staining intensity. The area of staining was scored as 0 (no tumor cells stained), 1 (< 25% of cells stained), 2 (25%–25% of cells stained). Staining intensity was graded as 0 (no staining), 1 (weak staining), 2 (moderate staining), 3 (strong staining).

**Total score was calculated by summing up the proportion score and intensity score of 2 different tissue microarray (TMA) cores. The area of staining was scored as 0 (0%–10% of the cells stained), 1 (11%–33% of the cells stained), 2 (34%–66% of the cells stained), and 3 (67%–100% of the cells stained). The staining intensity scored as 0 (no staining), 1 (weak staining), 2 (intermediate staining), and 3 (strong staining).
survival curves and P value with which to calculate HR with 95% CI.

2.2. Association between PD-L1 expression and overall survival

We investigated the correlation between PD-L1 expressions and OS in sarcoma patients. A total of 10 studies with 911 sarcoma patients were included in the analysis of overall survival. The combined HR of enrolled studies showed that PD-L1 expression was associated with poor OS (HR 1.35, 95% CI: 0.98–1.72, P = .000). Furthermore, a significant heterogeneity was detected among the studies with using random effects model for the analysis (I² = 48.0%, P = .044); therefore, we performed the subgroup analysis to clarify the potential sources (Fig. 2A).

**Figure 2.** Meta-analysis of the effect of PD-L1 expression on overall survival. The forest plot of hazard ratio for overall survival of patient with sarcoma (A), subgroup analysis stratified by the histological subtype (B), subgroup analysis stratified by the histological subtype (C), Subgroup analysis stratified by the cutoff (D). Subgroup analysis stratified by the target of PD-L1 assessment (E). PD-L1 = programmed cell death-ligands 1.
cutoff the heterogeneity was negligible ($I^2=0.0\%$, $P=.603$). Meanwhile, the pooled HR estimate for cutoff $\geq 5\%$ of tumor cells group was 1.44 ($95\%\; CI: 0.95–1.93$, $P=.000$) with medium heterogeneity ($I^2=64.1\%$, $P=.039$), as shown in Fig. 2D. To clarify the impact of a different PD-L1 assessment on the results, we conducted subgroup analysis stratified by the target of assessment. The pooled HR was 1.25 ($95\%\; CI: 0.85–1.66$, $P=.300$) for Protein, with high heterogeneity ($I^2=55.0\%$, $P=.300$), and 1.90 ($95\%\; CI: 0.94–2.85$, $P=.000$) for mRNA, with negligible heterogeneity ($I^2=0.0\%$, $P=.603$) (Fig. 2E).

### 2.4. Association between PD-L1 expression and event-free survival

Although 7 studies evaluated the association the PD-L1 expression with different effect size, including the events-free survival, disease-free survival, metastasis-free survival, and recurrence-free survival, the event was defined as local recurrence, distant metastasis, or death, which is accord with the DFS, MFS, RFS. Therefore, we combined the HRs as the effect size to evaluate survival outcome. The heterogeneity analysis showed low heterogeneity among the studies ($I^2=30.1\%$, $P=.198$), and the fixed-effect model was applied for HR. The pooled HR estimate was 1.70 ($95\%\; CI: 1.24–2.150$, $P=.000$) for the bone and soft-tissue sarcoma patients (Fig. 3). Overall, the meta-analysis showed a significant difference between positive and negative PD-L1 expression, revealed PD-L1 was a poor prognostic factor for events-free survival in sarcoma patients.

### 2.5. Association between PD-L1 expression and clinicopathological parameters

Seven studies assessed the association between PD-L1 expression and clinicopathological parameters. The clinical parameters analyzed included age, gender, tumor grade, tumor depth, tumor stage, tumor necrosis rate, tumor site, chemotherapy, radiotherapy, metastasis, recurrence. The pooled results of clinicopathological parameters are shown in Table 2. Furthermore, none of the clinicopathological parameters was found to be significantly correlated with PD-L1 expression. For the high heterogeneity group, a random effect model was used; meanwhile we also performed subgroup analysis, stratified by the histological subtype. The result still showed PD-L1 expression was not correlated with the clinicopathological parameters.

### Table 2

Relationship between PD-L1 protein expression and the clinicopathological feature.

| Clinicopathological features | Detection method | No. of study | Z (OR) | P value | $I^2$ | OR 95% CI | P value | Pool model |
|-----------------------------|------------------|-------------|--------|---------|------|-----------|---------|------------|
| Age (< 20 vs $\geq 20$)     | IHC              | 13,16,20    | 0.15   | .001    | 86.70% | 0.864 (0.557) | .878 | Random    |
| Gender (male vs female)     | IHC              | 16,17,20,21,22,23,24,25 | 0.77 | .762 | 0.109 (0.82–1.44) | .843 | Fix      |
| Tumor grade (low grade vs high grade) | IHC | 13,20,21,23,25 | 0.66 | .004 | 73.80% | 1.464 (0.472-4.542) | .51 | Random    |
| Tumor depth (superficial vs deep) | IHC | 13,20,25 | 0.45 | .045 | 67.70% | 1.251 (0.474-3.401) | .651 | Random    |
| Metastasis (yes vs no)      | IHC              | 16,17,20,21,23 | 1.54 | .000 | 58.50% | 2.801 (1.82–9.664) | .103 | Random    |
| Recurrence (yes vs no)      | IHC              | 16,17 | 0.57 | .648 | 0.128 (0.542–3.062) | .648 | Fix      |
| Tumor site (limb vs others) | IHC              | 21,22,25 | 1.24 | .105 | 55.60% | 1.705 (0.735–3.958) | .124 | Random    |
| Tumor stage (I/II VS III/IV) | IHC              | 13,20,22,23 | 0.08 | .024 | 68.20% | 0.943 (0.236–3.937) | .936 | Random    |
| Tumor necrosis rate $^*$ (high vs low) | IHC | 16,17,20 | 0.25 | .278 | 0.000 | 1.114 (0.478–2.946) | .802 | Fix      |
| Radiotherapy (yes vs no)    | IHC              | 13,21,22 | 0.77 | .001 | 84.90% | 1.781 (0.409–7.751) | .442 | Random    |
| Chemotherapy (yes vs no)    | IHC              | 13,16,21,22,23 | 0.02 | .999 | 0.000 | 1.176 (0.833–1.661) | .356 | Fix      |

$^*$ The high tumor necrosis rate for osteosarcoma (16, 17) means the tumor necrosis rate $\geq 90\%$, the high tumor necrosis rate for soft tissue sarcoma (20) means the tumor necrosis rate $\geq 50\%$. OR=odds ratio.
2.6. Publication bias and sensitivity analysis

By using Egger tests, there is no publication bias affecting the hazard ratios for overall survival and events-free survival was present in the eligible studies. The P values for these tests were .679 and .163, respectively. At the same time, no publication biases were observed based on the visual evaluation of the Begg funnel plot (Fig. 4A, B).

Sensitivity analysis, by omitting 1 study at a time, was performed to detect the stability of the results. The results shown in Fig. 5A, B demonstrated that no unique study significantly affected the pooled HRs of OS and EFS, indicating that the results of the present meta-analysis are stable and credible.

3. Discussion

Bone and soft tissue sarcoma are a rare heterogeneous neoplasms of mesenchymal derived, accounting for approximately 1% of all human malignance.[27] Although the overall survival rate has been raised with the development of effective chemotherapy and surgical techniques, the cures for sarcoma patients by using the traditional strategy were stagnant for decades.[28–30] However the local recurrence and metastasis are still common and the average survival of these patients is poorer.[31] In the contrast, the immunotherapy has achieved tremendous success in solid tumors and been considered the foremost method in individualized medicine.[5,32] Understanding how immunology mechanism of action relates to sarcomas may reveal the potential for immunotherapy and to develop new effect treatment method for sarcoma.

PD-1 and its ligands, PD-L1 and PD-L2 overexpressed within the tumor microenvironment, can inhibit T-cell activation and proliferation and negatively regulate the immune response through various pathways in antitumor immunity, thereby losing its killing effect on tumor cells and protecting tumor cells from the host immunologic surveillance system.[33] Recently, the association between PD-L1 expressions and various solid tumors has been studied by numerous researches and the results have shown that expression of PD-L1 significantly correlates with poor prognosis.[9,34–36] However, its expression and impact on the survival outcome of patients with bone and soft tissue sarcoma remains inconsistent and conflicting. Multiple studies have shown that positive PD-L1 was associated with significantly poor prognosis,[13,14,17,20,21,26] but other studies could not confirm this finding. To achieve a reasonable conclusion, we conducted this meta-analysis and aimed to assess the correlation between PD-L1 expression and the prognosis of bone and soft tissue sarcoma patients.

According to the result of literature retrieval, our analysis combined14 studies (including 15 cohorts) with 1451 patients. The results reveal that PD-L1 expression is a negative prognostic factor in bone and soft tissue sarcoma with statistical significances for OS (HR = 1.35, 95% CI: 0.98 – 1.72), and EFS (HR = 1.70, 95% CI: 1.24 – 2.150). Due to the high heterogeneity among the overall survival of sarcoma patients, we preformed
subgroup analysis according to different histological subtypes, ethnicities, target of PD-L1 assessment, and cut-off. Interestingly, for tumor histological subtype, the analysis results have shown a significant association between PD-L1 expression and the poor overall survival of patient with osteosarcoma, soft tissue sarcoma, with no significant heterogeneity ($I^2 = 0.0\%, P = .486$; $I^2 = 42.8\%, P = .120$), respectively. Therefore, we consider that the different histological subtype might be a factor that explains the heterogeneity. In the enrolled studies, the different cutoff and target of PD-L1 assessment were used to detect the expression of PD-L1, while the pooled HR for protein and cutoff $\geq 5\%$, $< 5\%$ group show high heterogeneity ($I^2 = 55.0\%, P = .030$ and medium heterogeneity ($I^2 = 64.1\%, P = .038$; $I^2 = 44.5\%, P = .144$), respectively. We presume that even IHC is the common method used by including studies, PD-L1 positivity was assessed by using different antibodies, the sensitivities of the antibodies and the experimental procedures, different cutoff might be the factor which contributes to the heterogeneity.

According to the result of included studies in the meta-analysis, the result was consistent with multiple studies, however, expression of PD-L1 was a favorable prognostic factor and improved the survival in 1 study. These inconsistent and conflicting results have several possible explanations: different histology type of tumor was selected to be investigated and different sample size from different areas of study which suggested that result of study might be debatable. Meanwhile, comparisons of different studies reporting PD-L1 expression are unconformity by the use of different methodologies, different thresholds, different antibodies. Additionally, tumor cells express PD-L1 by antitumor immune response and oncogene-driven mechanisms. The former is induced by cytokines and dependent on the presence of T lymphocytes infiltration. The latter does not depend on the presence of T lymphocytes infiltration and multiple mechanisms can lead to PD-L1 expression, we hypothesize that these inconsistent results might be partially due to the impact of PD-L1 expression on the different populations of T lymphocytes infiltration or active immune responses and different PD-L1 regulation at both transcription and translational levels. Further investigations will be required to clarify the underlying mechanism.

In published data, nearly all the clinicopathological features, including age, gender, tumor grade, tumor depth, tumor stage, tumor necrosis rate, tumor site, chemotherapy, radiotherapy, metastasis, recurrence, have been demonstrated associated with PD-L1 expression in sarcoma. However most of them were not strongly confirmed by other studies. Only 1 study showed PD-L1 expression was associated with age. Meanwhile, PD-L1 expression was found to be associated with tumor necrosis rate also by only 1 study. With regards to tumor grade, tumor stage, and tumor depth, significant results were both reported in 2 studies. Besides, PD-L1 expression was found to be correlated with tumor metastasis in 5 studies. However, when we combined the data together, none of the clinicopathological features mentioned above was associated with PD-L1 expression. The discordances among previous and current analysis may result from inadequate sample size, heterogeneous study population, and variable definitions of PD-L1 expression and different methodologies.

Thus, studies with larger sample size and standardized quantitative assays were still needed to assess the correlation between PD-L1 expression and clinicopathological parameters. Although we made an effort to perform a comprehensive analysis, but there were some limitations to our analysis. First, significant heterogeneity was observed among studies, which will influence the conclusions of this study. To minimize the effect of the heterogeneity, a random effect model and subgroup analysis were applied for high heterogeneity group. Second, within each study, different methods of PD-L1 measurement were used to detect the expression of PD-L1. Despite that the common method was IHC, PD-L1 positivity was assessed using different antibodies, as the sensitivities of the antibodies, the experimental procedures, and cutoff are varied. Meanwhile, the immunohistochemical reagents, the scoring method, the cut-off values used to determine PD-L1 positivity also varied among the studies included in the analysis. Thus, these variances of different methodology may result in heterogeneity among the including studies of this meta-analysis and may lead to a potential bias. Third, the sample sizes of the studies eligible in the analysis were relatively small. However, the results of the sensitivity analysis results still remain stable and creditable after the sequential exclusion of each study. Fourth, not all of the HR with 95% CI was directly extracted from the studies, so if this was impracticable we had to extrapolate the HR via Kaplan–Meier curves or P values. The estimation might be less reliable than those reported directly. Finally, publication bias should be concerned, our meta-analysis was limited to articles published in English. Additionally, certain studies with negative results or some useless results may not be published in journals, which may result in publication bias. To minimize the effect, we conduct comprehensive literature searches as completely as possible by using PubMed, Embase, Web of Science, and the Cochrane Library. But no statistical significant publication bias was found in this meta-analysis. Nevertheless, it should be noted that we excluded the case reports, reviews, letter, and conference abstracts as it did not contain sufficient data for aggregation. In general, these above limitations may bring potential source of publication bias to the current meta-analysis. Therefore, the conclusion and results of the current meta-analysis should be interpreted with caution and need to be validated by more well-designed prospective studies with appropriate multivariate analyses.

4. Conclusion

In conclusion, this meta-analysis demonstrates that PD-L1 expression is significantly correlated with poor OS and EFS for patients with sarcoma, and PD-L1 expression may be a useful predictive factor of prognosis for bone and soft tissue sarcoma. This information may be beneficial to clinicians to choose individual treatment methods by anti-PD-1/PD-L1 therapy. Future adequately designed clinical studies with uniform assessment assay are needed to verify the role of PD-L1 expression in sarcoma.

5. Materials and methods

5.1. Literature search strategy

A comprehensive literature search was performed in the PubMed, Embase, Web of Science, and Cochrane databases with no language restrictions. Articles published before August 2017 were included in the meta-analysis. The following keywords were used as both text words and Medical Search Headings (Mesh terms) and were combined by using Boolean operators for the relevant literature: (“Sarcoma”[Mesh] OR “Sarcoma”∗ OR “Soft tissue sarcoma”∗ OR “Epithelioid Sarcoma”∗ OR “Spindle Cell Sarcoma”∗ OR “Osteosarcoma”∗ OR “Osteosarcoma...
Tumor*” OR “Osteogenic Sarcoma*” OR “Chondrosarcoma*” OR “Ewing sarcoma*” OR “Ewing’s Tumor*” OR “Carcinosarcoma*” OR “Leiomyosarcoma*” OR “Angiosarcoma*” OR “Desmoplastic Small Round-Cell Tumor*” OR “Hemangiosarcoma*” OR “Lymphangiosarcoma*” OR “Malignant Lymphangiendothelioma*” OR “Myosarcoma*” OR “Fibrosarcoma*” OR “Synovial sarcoma*” OR “Malignant Fibrohistiocytic Tumor*” OR “Malignant Fibrous Histiocytomas*” OR “Liposarcoma*” OR “Dedifferentiated Liposarcoma*” OR “Pleomorphic Liposarcoma*”) AND (“PD-L1” OR “B7-H1” OR “CD274” OR “programmed cell death 1 ligand 1 protein” OR “CD274 Antigen” OR “B7H1 Antigen” OR “PD-L1 costimulatory protein” OR “B7-H1 Immune Costimulatory Protein” OR “PD-L1 Costimulatory Protein”).

5.2. Eligibility criteria and quality assessment

The eligible studies for the analysis had to be in accordance with the following criteria: studies whose entire populations comprised patients with histologically confirmed bone and soft tissue sarcoma; studies providing sufficient data regarding the correlation between PD-L1 and clinicopathological features and overall survival (OS); studies that revealed the prognosis provided the sufficient information to estimate hazard ratio (HR) about survival outcome. The language of literature was restricted to English. Studies were excluded from the analysis if they meet the following: reported overlapping patients; nonclinical research; reviews, case reports, letters and articles from conferences; with insufficient information. When the same patient source was included in different publications, the most recent and largest study was included in analysis. Two independent reviewers (CZ and WY) verified the included studies met the above criteria for subsequent analysis. Since all the studies included were cohort studies, the quality of each study was evaluated using Newcastle–Ottawa Scale (NOS) (www.ohri.ca/programs/clinical_epidemiology/oxford.asp). Based on the assessment of each study in selection, comparability, and exposure, a score up to 9 points was appointed. The quality assessment was carried out independently by 2 reviewers (CZ and WY). The disagreement was resolved by consensus (CZ, WY, and SZ).

5.3. Data extraction

All relevant data of the eligible articles were extracted by 2 independent researchers (CZ and WY), and any disagreements were resolved by consultation. The following data was extracted: basic information of each article including first author, year of publication, country, number of patients, follow-up duration, and study design; data of patient and tumor including age, gender, tumor grade, tumor depth, tumor stage, tumor necrosis rate, tumor site, chemotherapy, radiotherapy, metastasis, recurrence; prognosis outcome measures including overall survival, disease-free survival, events-free survival, recurrence-free survival, metastasis-free survival, Kaplan–Meier curves and P values; other data including the assay methods and PD-L1 positivity expression cut-off value.

5.4. Statistical analyses

To evaluate the prognostic significance of PD-L1 expression, HRs and theirs 95% CIs were used to determine the association between PD-L1 expression and survival, and pooled ORs and its 95% CIs were used to assess the correlation between PD-L1 and clinicopathological parameters of sarcoma patients. When the HRs were given explicitly in the article, we used the original data, or calculated the HRs with 95% CIs from available survival data in the articles or from Kaplan–Meier curves by using the methods reported by Tierney et al. [90]

Statistical heterogeneity among each study was evaluated by using Cochran Q test and Higgins I² statistic. If \( P > 0.1 \) and \( I^2 < 50\% \), it indicated a significant heterogeneity between studies and a random effects model was selected to combine the data in these cases. Otherwise, the heterogeneity was not significant and a fixed effects model was used. Meanwhile, the subgroup analysis was further performed to explore the heterogeneity source. Publication bias was assessed by Egger test and Begg test. All statistical analyses were performed using STATA version 12.0 (Stata Corp, College Station, TX). \( P < 0.05 \) were considered statistically significant.

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