Immune checkpoint inhibitors for esophageal squamous cell carcinoma: a narrative review

Wencheng Zhang, Ping Wang, Qingsong Pang

Department of Radiation Oncology, Tianjin Medical University Cancer Institute and Hospital/National Clinical Research Center for Cancer, Tianjin, China

Contributions: (I) Conception and design: Q Pang; (II) Administrative support: P Wang; (III) Provision of study materials or patients: W Zhang; (IV) Collection and assembly of data: W Zhang; (V) Data analysis and interpretation: W Zhang; (VI) Manuscript writing: All authors; (VII) Final approval of manuscript: All authors.

Correspondence to: Qingsong Pang; Ping Wang. Department of Radiation Oncology, Tianjin Medical University Cancer Institute and Hospital, Huanhu West Road, Hexi District, Tianjin, China. Email: pangqingsong@tjmuch.com; wangping@tjmuch.com.

Abstract: Esophageal cancer (EC) has the seventh highest incidence and the sixth highest mortality rate of any type of cancer worldwide. In China, esophageal squamous cell carcinoma (ESCC) accounts for more than 95% of EC patients. The main treatment for EC patients is surgery and/or chemoradiotherapy (CRT). A large proportion of EC patients are already at an advanced stage of the disease by the time they are diagnosed. In these cases, CRT is left as the only treatment choice, and the treatment outcome is poor. Immune checkpoint inhibitors (ICIs) can improve clinical response and patient survival of patient with many types of tumors through reactivating antitumor immune response. The study of ICIs in ESCC is relative delayed compared with that in other solid tumors. Recent results from clinical trials have demonstrated the safety and efficacy of ICIs either alone or combined with chemotherapy or chemoradiotherapy in ESCC patients. Accumulated evidences also have shown the improved treatment outcome was associated with PD-L1 expression, tumor DNA instability-induced tumor mutational burden (TMB), and drawing lymphocytes into the tumor. Based on these findings, ICIs combined with CRT or radiotherapy (RT) are the focus of ongoing studies. This review will summarize the recent progress in this field, especially the mechanism of ICIs used in ESCC, their clinical efficacy and toxicities, and potential biomarkers.

Keywords: Esophageal squamous cell carcinoma (ESCC); immune checkpoint inhibitor (ICI); immunotherapy; chemotherapy; radiotherapy (RT); biomarker

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Introduction

In 2018, there were approximately 570,000 new cases of esophageal cancer (EC) and around 510,000 related deaths globally, placing it seventh and sixth among all cancer types in terms of morbidity and mortality, respectively (1). China accounts for half of EC cases worldwide. EC comprises two histological subtypes, esophageal adenocarcinoma (EAC) and esophageal squamous cell carcinoma (ESCC), which each have distinct pathological features and molecular characteristics (2). In contrast with Europe and the United States, where EAC is more common, 95% of EC cases in China are ESCC. Surgery remains the mainstay for treatment of ESCC at early stage. Several randomized controlled trials demonstrated that preoperative chemoradiotherapy (CRT) compared with surgery alone increases the overall survival (OS), but the treatment-related adverse effects (TRAEs) raised either (3,4). In China, 60% to 70% of patients have already developed locally advanced or advanced disease at the time of diagnosis (5). For these patients, concurrent CRT (CCRT) is the only choice. But the treatment outcome remains poor. Although the
antibodies that target epidermal growth factor receptors have been investigated in ESCC, the results from clinical trials did not indicate positive results (6). As a result, new and effective strategies for advanced ESCC are urgently needed.

T cells plays an important role in immune defense against cancer. T-cell activation is strictly regulated by stimulatory and inhibitory ligand-receptor interaction between T cells, dendritic cells, macrophages and tumor cells in the tumor microenvironment. These ligand-receptor pairs that negatively regulates T-cell activation are named “immune checkpoints”. A growing number of studies have demonstrated that immune checkpoint inhibitors (ICIs), such as anti-CTLA-4 mAb (ipilimumab) and anti-PD1 mAb (nivolumab and pembrolizumab), can promote tumor shrinkage and improve overall survival in a variety of tumors (Figure 1). Since ipilimumab was approved firstly by FDA for the treatment of metastatic melanoma in 2011, additional ICIs that all target the PD-1/PD-L1 axis have been approved for the treatment of a broad range of tumor types. Research on ICIs in ESCC started later, while several ongoing clinical trials featuring anti-PD-1 mAb have shown promising results. The development of ICIs has been explored (7). This review focused on clinical application of ICIs for treating ESCC, including the feasibility and efficiency of mono or combined treatments of ICIs, the safety and TRAEs as well as biomarkers that were studied as hotspots in recent years. We present the following article in

Figure 1  T cell inhibition and T cell activation by checkpoint inhibition. (A) The binding of CTLA-4 and CD80/CD86 that were expressed on antigen presenting cells blocked the downstream transduction of activation signaling resulting from the recognition of T-cell receptor and MHC-antigen complex. CTLA-4 inhibitors that targeted CTLA-4 prevented the binding of CTLA-4 from CD80/CD86 and restored the initial activation of T cells. (B) The interaction of PD-1 on activated T cells and its ligands PD-L1/PD-L2 on tumor cells, antigen presenting cells and regulatory T cells inhibited the continuous activation of T cells. PD-1/PD-L1 inhibitors that targeted the PD-1/PD-L1/2 axes could maintain the continuous activation of T cells.
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**ICIs required for ESCC treatment**

Heterogeneous factors in the tumor microenvironment limit antitumor immunity and support tumor evasion. The function of antitumor effector T cells is inhibited by the interaction between checkpoint molecules and their ligands expressed on tumor cells, stromal cells, and immune cells, which ultimately leads to immune escape (8,9). In ESCC, the checkpoint ligand PD-L1 has been investigated most extensively. PD-L1 expression on tumor cells is regulated by both intrinsic and exogenous factors (10) (Figure 2). Ohigashi et al., who evaluated PD-L1 or PD-L2 gene expression in 18 (43.9%) of 41 ESCC tumor samples using real-time PCR method, showed for the first time that PD-L1 was a novel prognostic marker for human EC (11). Meanwhile, in a meta-analysis involving 1,350 ESCC patients from China and Japan, PD-L1 overexpression was found in 559 patients (41.4%) and indicated to be a poor prognostic biomarker for ESCC (12). Immunohistochemistry (IHC) was the only method for evaluating PD-L1 expression used in this study (12). Tsutsumi et al. also reported an association between PD-L1 expression and worse rates for OS and relapse-free survival (RFS) (13). Patients with positive PD-L1 expression have significantly lower pathological complete response rates (13% versus 32%; P=0.036) than those with negative PD-L1 expression. In addition, Huang et al. discovered that PD-L1 expression influenced the effect of neoadjuvant CRT in ESCC patients, finding PD-L1 expression to be an independent predictor of poor CRT response and worse treatment outcome (14).

Tumor mutational burden (TMB) represents the number of genetic changes in a tumor. Tumors with a high TMB are likely to express high levels of neoantigens, and therefore could potentially possess high capability to attract T cells that participate in antitumor immune response. Several studies have shown high TMB in ESCC (15,16). Yarchoan et al. observed a significant correlation between TMB and the objective response rate (ORR) (17). Microsatellite instability (MSI) is caused by the failure of the DNA mismatch repair (MMR) system, which is usually responsible for repairing DNA. High MSI (MSI-H) typically correlates with increased neoantigen burden, resulting in a much greater response to ICIs. The anti-PD-1 mAb pembrolizumab (Keytruda) is the first FDA-approved treatment based on MSI-H or MMR deficiency, regardless of cancer types. Studies on MSI have shown a much lower level of MSI-H in ESCC than in gastric cancer (2). The relatively high levels of PD-L1 expression and TMB in ESCC, as well as MSI-H in a proportion of patients, indicate that patients with ESCC might be susceptible to ICI treatment.

Tumor-infiltrating lymphocytes (TILs), which influence clinical response and survival, are important components in tumor microenvironment. With increased checkpoint expression and reduced antitumor function, CD4+ helper T cells and CD8+ effector T cells in tumors become exhausted in the inhibitory immune microenvironment. In contrast, inhibitory TILs, such as the regulatory T cells (Tregs) and

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**Figure 2** PD-1/PD-L1 expression in esophageal squamous carcinoma cell. PD-1 expressed on the membrane of tumor-infiltrating lymphocytes (A) and PD-L1 expressed on the membrane of tumor cells (B) were assessed by immunochemistry assay (200x).
tumor-associated macrophages (TAMs), are activated by the elevated production of IL10, IL6, transforming growth factor-β (TGF-β), and expression of checkpoint molecules. Meanwhile, the density and location of TILs also affect their role in tumors. Tumors can be classified according to the characteristics of TILs into four categories: hot, altered-excluded, altered-immunosuppressed, and cold. Hot tumors, which are also defined as inflamed tumors, have high levels of TILs. In altered-excluded tumors, TILs are excluded around the tumor margins, while in immunosuppressed tumors, TILs are either scattered in or surround the tumor. Owing to the immunosuppressive microenvironment, the number of TILs in immunosuppressed tumors is much lower than that in altered-excluded tumors. Finally, cold tumors, also known as non-inflamed tumors, are devoid of TILs. Although no study has identified tumor immune categories in EC, it has been reported that a high density of CD8 cells was significantly correlated with more favorable OS and RFS in patients who received post-CRT (18). Similarly, we found that high levels of TILs induced by CCRT were associated with improved clinical response and survival (19). Furthermore, the effect of intratumoral location of TILs on the survival of ESCC patients varies. High numbers of CD8+ and PD-1+ TILs only in the surface and center of the tumors were related to improved OS (20). Optimal therapeutic strategies that target the different tumor categories should be provided. Of these four tumor categories, hot tumors may have higher susceptibility to PD-1/PD-L1 blockade. Meanwhile, CRT and RT combined with immunotherapy might be optimal for the other tumor categories in inducing and activating antitumor immune response and attracting more functional T cells into tumors.

**The use of ICI s in EC (Table 1)**

**Pembrolizumab**

In a phase Ib study from KEYNOTE-028 that explored the efficacy and safety of pembrolizumab (anti-PD-1 mAb) in late-line treatment (87% patients had received ≥2 lines of treatment) for all histological types of EC (78% ESCC, 22% EAC), Twenty-three patients had PD-L1-positive tumors. The mOS was 7 months (21). The phase II trial KEYNOTE-180 investigated pembrolizumab monotherapy in late-line (≥3) therapy for patients with PD-L1-positive tumor. ESCC and EAC were both included. The mOS and median PFS (mPFS) were 5.8 and 2 months, respectively. Subgroup analysis showed that the ORR and mOS were better in patients with ESCC (22). In the phase III trial from KEYNOTE-181, pembrolizumab used in the second or later line of therapy did not have significant survival benefit compared with chemotherapy. However, in the subgroup of patients with PD-L1 combined positive score (CPS) (PD-L1 expression on tumors and immune cells were both included) ≥10, pembrolizumab treatment achieved a longer mOS than chemotherapy (9.3 vs. 6.7 months), and the ORR was also improved (21.5% and 6.1%) in this subgroup (23).

**Nivolumab**

ATTRACTION-1, a Japanese phase II study, investigated the effect of nivolumab (anti-PD-1 mAb) on ESCC. The ORR was 17%, and the mOS and mPFS were 10.8 and 1.5 months, respectively (24). ATTRACTION-03 (NCT02569242) enrolled patients with unresectable advanced or recurrent ESCC. The mOS was significantly improved in the nivolumab group compared with that in the chemotherapy group (10.9 vs. 8.4 months) (25). Meanwhile, the CheckMate-032 study (NCT01928394) assessed the efficacy and safety of nivolumab and nivolumab plus ipilimumab in three groups of EC patients (nivolumab 3 mg/kg; nivolumab 1 mg/kg plus ipilimumab 3 mg/kg; and nivolumab 3 mg/kg plus ipilimumab 1 mg/kg). The ORR was 12%, 24%, and 8% (95% CI, 2% to 19%) in the three groups, respectively (26). The RAMONA study, a multicenter, open-label phase II trial, has the primary objective of demonstrating the significant survival benefit of nivolumab and ipilimumab in older patients with advanced ESCC based on a comparison with historical data of standard chemotherapy; the study is ongoing (27).

**Camrelizumab**

One trial (NCT02742935), a dose-escalating phase I study, evaluated the efficacy and safety of camrelizumab (also name SHR-1210, humanized anti-PD1 IgG4 antibody developed by Jiangsu Hengrui Medicine Co. Ltd, China) as a second or later-line treatment for ESCC. The dose was given at 60, 200, and 400 mg every 2 weeks. The ORR was 33.3% and the mPFS was 3.6 months (28). The ESCORT (NCT03099382) study to assess the efficacy and safety of camrelizumab vs. chemotherapy in patients with advanced or metastatic ESCC after the failure of first-line standard therapy, was the first randomized phase III study conducted
| Trial       | Agent          | Histology          | Arms   | N   | Phase   | Line          | Number | PD-L1 positive | ORR  | mPFS (m) | mOS (m) | 1-year PFS | 1-year OS |
|-------------|----------------|--------------------|--------|-----|---------|---------------|--------|----------------|------|----------|---------|------------|------------|
| KEYNOTE-028 | Pembrolizumab  | SCC                | Single | 18  | phase IB | 2 or more     | 23     | 100%           | 28% | 1.8      | 7.0     | 22%        | 40%        |
|             |                | AC                 |        |     |         |               |        |                |      |          |          |            |            |
| KEYNOTE-180 | Pembrolizumab  | SCC                | Single | 63  | phase 2 | 2 or more     | 121    | 47.9%          | 14.3%| 2.1      | 6.8     | 12% (9 m) | 34%        |
|             |                | AC                 |        |     |         |               |        |                | 5.2%| 1.9      | 3.9     | 7% (9 m)  | 22%        |
| ATTRACTION-1| Nivolumab      | SCC                | Single | 64  | phase 2 | 2 or more     | 64     | NA             | 22% | 2.3      | 10.8    | 11%        | 43%        |
| NCT02915432| Toripalimab    | SCC                | Single | 56  | phase 1b/2 | 2 or more  | 56     | 21.4%          | 23.5%| NA       | NA      | NA         | NA         |
| NCT02742935| Camrelizumab   | SCC                | Single | 30  | phase 1 | 2 or more     | 30     | 62.5%          | 33.3%| 3.6      | NA      | 31.5% (6 m)| NA         |
| KEYNOTE-181 | Pembrolizumab  | SCC/AC             | Double | 314 | Phase 3 | 2 or more     | 628    | 34.1%          | NA   | NA       | 7.1     | NA         | 18%        |
|             | Chemotherapy   | SCC/AC             |        | 314 |         |               |        | 36.6%          | NA   | NA       | 7.1     | NA         | 10%        |
| KEYNOTE-181 | Pembrolizumab  | SCC/AC             | Double | 107 | Phase 3 | 2 or more     | 222    | 100%           | NA   | NA       | 9.3     | NA         | 43%        |
| (CPS ≥10)   | Chemotherapy   | SCC/AC             |        | 115 |         |               |        | 100%           | NA   | NA       | 6.7     | NA         | 20%        |
| KEYNOTE-181 | Pembrolizumab  | SCC or AC          | Double | 60/2| Phase 3 | 2 or more     | 62     | 40.3%          | NA   | NA       | 8.4     | NA         | 36%        |
|             | Chemotherapy   | SCC or AC          |        | 59/2|         |               |        | 47.5%          | NA   | NA       | 5.6     | NA         | 17%        |
| ESCORT      | Camrelizumab   | SCC                | Double | 228 | Phase 3 | 2 or more     | 448    | 40.8%          | 20.2%| 1.9      | 8.3     | 10%        | 33.7%      |
|             | Chemotherapy   | SCC                |        | 220 |         |               |        | 44.5%          | 6.4% | 1.9      | 6.2     | NA         | 22.3%      |
| ATTRAC-     | Nivolumab      | SCC                | Double | 210 | Phase 3 | 2 or more     | 419    | 48%            | 19%  | 1.7      | 10.9    | 12%        | 47%        |
| TION-03     | Chemotherapy   | SCC                |        | 209 |         |               |        | 49%            | 22%  | 3.4      | 8.4     | 7%         | 34%        |

ESCC, esophageal squamous cell carcinoma; AC, adenocarcinoma; N, number of patients; ORR, objective response rates; mPFS, median progression-free survival; mOS, median overall survival; NA, not available.
in a Chinese population. The results revealed improved mOS and a 31% reduction in the risk of progression or death after camrelizumab treatment compared with after chemotherapy, as well as superior ORR (20.2% vs. 6.4%) and duration of response (7.4 vs. 3.4 months).

**Toripalimab**

A phase Ib/II trial (NCT02915432) was conducted to evaluate the efficacy and safety of toripalimab (humanized anti-PD-1 mAb developed by Shanghai Junshi Bioscience) in refractory/metastatic ESCC. Of 34 patients involved in the trial, 8 achieved partial response, with an ORR of 23.5% (29).

**ICIs combined with chemotherapy in ESCC**

Two mechanisms are involved in chemotherapy-induced immune response. First, chemotherapeutic drugs influence the immune effect of tumor cells. The capacity of cyclophosphamide, gemcitabine, platinum, and paclitaxel to enhance the antigens of tumor cells has been shown (30). Paclitaxel, cisplatin, and doxorubicin could increase the immunogenicity of tumor cells by promoting immunogenic cell death of tumor cells and the expression and release of immunogenic substances from tumor cells, as well as by enhancing the immune effect sensitivity of tumor cells (31). Second, chemotherapeutic drugs can influence immune cells. Cytotoxic T lymphocytes are activated after treatment with Taxol, doxorubicin, or cisplatin. Taxol, gemcitabine, and 5-fluorouracil can eliminate immunosuppressive cells. Changes in the tumor microenvironment, such as an increase in the release of tumor antigens, promote antigen cross-presentation, and thus accelerate anti-tumor immune response (31). The effects of chemotherapy on tumor cells and immune cells reciprocate, and re-shape the tumor immune response.

Several studies have demonstrated that immunotherapy combined with chemotherapy improved ORR and survival in multiple tumors and that the TRAEs are well tolerated. Since neoadjuvant chemotherapy has been shown to increase PD-L1 expression and CD8+ TILs in ESCC (32), chemotherapy combined with PD-1/PD-L1 blockade might achieve a potential clinical outcome. Although no reports currently exist, two randomized, double-blind, placebo-controlled, multicenter phase III clinical trials are under way. The first of these trials, KEYNOTE-590 (NCT03189719), will evaluate the efficacy and safety of cisplatin and 5-fluorouracil plus pembrolizumab vs. cisplatin and 5-fluorouracil plus placebo in patients with previously untreated advanced E/EGJ carcinoma (33). The second, SHR-1210-III-306 (NCT03691090), will compare the efficacy and safety of SHR-1210 + paclitaxel + cisplatin vs. placebo + paclitaxel + cisplatin as first-line therapy for patients in China with advanced EC.

**ICIs combined with RT in ESCC**

The combination of ICIs with RT has drawn an increasing level of attention in recent years. RT boosts *in situ* vaccination by inducing immunogenic death in radiated tumor cells. RT reprograms the tumor immune microenvironment through the release of inflammatory cytokines and chemokines, which contributes to the recruitment of effector T cells in tumors and converts the tumors into “inflamed” tissues which are susceptible to T-cell attack. Our data demonstrated an elevated expression of PD-L1 on ESCC cell lines after radiation (10). Therefore, when immune response was promoted by RT-induced tumor destruction, PD-L1-expressing tumor cells were also selected. For patients who received esophagectomy after neoadjuvant CCRT, PD-L1 staining was linked with lower complete pathologic response rate [16% (3/18) in PD-L1+ patients vs. 31% (9/29) in PD-L1- patients] (34). Preclinical and early clinical data have revealed the synergistic effect of RT and PD-1/CTLA-4 blockade, which has led to an increase in interest in the combined use of these two treatment strategies (35). In our phase Ib clinical trial to evaluate the safety and effect of RT combined with SHR-1210 as a first-line treatment in locally advanced ESCC patients who were intolerant to CCRT, 19 patients were finally enrolled. The ORR was 73%, and the mPFS and mOS were 11.7 and 16.9 months, respectively. At 12 months, the rates of OS and PFS were 63.2% and 47.4%, respectively. The most common type of toxicity was cutaneous capillary hemangioma (15 in grade 1 and 2 in grade 3). Grade 3 TRAEs occurred in 9 patients experienced (31). One patient experienced grade 4 lymphopenia, which was not treated (36).

**ICIs combined with CRT in ESCC**

A number of clinical trials are being conducted to investigate the efficacy of immunotherapy combined with neoadjuvant CRT in esophageal and gastric cancer or gastroesophageal junction adenocarcinoma (NCT03064490, NCT02730546,
NCT02735239, NCT02962063, NCT03044613 and NCT03278626). The primary outcomes from these phase I/II clinical trials are safety and/or the rate of pathologic complete response as determined at the time of surgery. In addition to these trials, others are incorporating immunotherapy into the definitive, combined modality management of EC (NCT03377400 and NCT03544736).

In a phase II trial (NCTNCT02844075), patients with stage Ib to III ESCC received pembrolizumab combined with neoadjuvant CCRT over 5 weeks, followed by surgery, and for another two years after surgery. Of the 28 patients enrolled, 26 patients received esophagectomy, while 2 patients did not undergo surgery due to death (hematemesis) and the withdrawal of consent. After surgery, two patients died from acute lung injury. The pathologic complete response (pCR) in the primary tumor was 46.1%. The 6- and 12-month OS rates were 89.3% and 82.1%, respectively. The most common TRAEs in the neoadjuvant and adjuvant periods were neutropenia (50.0%) and liver enzyme elevation (30.8%), respectively (37). Another phase II clinical trial will evaluate the safety and efficacy of atezolizumab (anti-PD-L1 mAb) monotherapy following definitive CRT in patients with unresectable locally advanced ESCC. Within 4 weeks after CRT, the patients will receive atezolizumab every 3 weeks for 12 months or until disease progression. The trial will enroll a total of 50 patients including 40 with locally advanced primary ESCC and 10 with postoperative loco-regionally recurrent ESCC (38). Other than ESCC-focused studies, there is a phase II trial (NCT03087864) that is investigating neoadjuvant CRT combined with atezolizumab in resectable EAC. Of the 39 patients enrolled, 31 patients have completed neoadjuvant treatment and 8 patients are still receiving treatment. A pCR has been observed in 39% of patients, which is promising compared to the 23% in the CROSS study in which neoadjuvant CRT was applied. Treatment-related immune AEs are manageable (39). The preliminary results from ongoing clinical trials have demonstrated that the addition of PD-1/PD-L1 blockade to preoperative CRT or definite CCRT has promising efficacy with manageable toxicity in EC. Based on the results, further investigation is warranted in a phase III clinical trial.

**Other combination treatment**

A single-center phase II study evaluated the efficacy and safety of camrelizumab plus apatinib, a tyrosine kinase inhibitor of vascular endothelial growth factor receptor-2 (VEGFR2-TKI), in combination with liposomal paclitaxel and nedaplatin in the first-line treatment of patients with ESCC. Response evaluation was available in 26 patients of 29 patients included. The ORR and DCR were 73.1% (19/26) and 96.2% (25/26), respectively. The most common grade 3/4 adverse events were leucopenia (21/29, 72.4%) and neutropenia (15/29, 51.7%). The preliminary result indicated that ICIs plus VEGFR2-TKI in combination with chemotherapy might provide new treatment option for patients with unresectable locally advanced or metastatic ESCC (40).

**Treatment-related adverse effects**

A meta-analysis of 18,610 patients treated with PD-1 and PD-L1 inhibitors revealed that 66.0% patients developed at least 1 AE of any grade, and 14.0% patients developed at least 1 AE of grade 3 or higher severity (41). Different tumors appeared to have various TRAEs (21,42). Several clinical trials have reported the TRAEs in ESCC patients treated with PD-1 or PD-L1 inhibitors (Table 2). TRAEs of any grade were less likely with pembrolizumab (39–65%) and nivolumab (60–65%) than with camrelizumab (83.3–94.3%). The occurrence of grade 3–5 TRAEs was similar among pembrolizumab (12.4–18.2%), nivolumab (17–18%), and camrelizumab (10–19.3%), which was lower than chemotherapy (39.5–63%). The ongoing clinical trials have indicated that combining ICIs and chemotherapy or CCRT does not increase the occurrence of TRAEs compared with chemotherapy or CCRT. TRAEs include diarrhea, decreased appetite, constipation, fatigue, and immune-related (irAEs), such as rash, hyperthyroidism, hypothyroidism, pneumonitis, and colitis. irAEs can be controlled by suspending the treatment or with corticosteroid treatment. It should be noted that there is a high incidence (83.3%) of reactive capillary hemangiomas in camrelizumab which are mostly of grade 1–2 and manageable with local therapy or observation. In order to prevent TRAEs, it is important to exclude patients who had serious baseline disease, such as in lung and digestive tract, before ICIs application. These patients are more susceptible to TRAEs. After ICIs application, the core principles of treating irAEs are early prevention and diagnosis as early as possible, and performing proper management. It is also necessary to monitor for the recurrence of irAEs when ICI treatment is resumed.
Biomarkers for predicting the efficacy of ICIs in ESCC (Table 3)

The reported studies indicated the ORR of 10–30% in ESCC patients treated with ICIs, and there is still a large proportion of patients who do not benefit from ICI treatment. There is an urgent need for potential predictive biomarkers to select the most suitable patients, optimize therapeutic strategies, and monitor the clinical response. Unlike the precise targets and definite mechanism in targeted therapy, much more complicated factors arising from tumor cells and the tumor immune microenvironment harmonize and contribute to the clinical outcome of ICI-based treatment. A series of immune-related biomarkers, such as PD-L1 expression, TMB, MSI, and TILs, have been observed as valuable predictive candidates in ICI-based treatment for ESCC (43).

**PD-L1**

An increasing level of evidence has shown that PD-L1 expression in tumor cells is potential biomarker for predicting the therapeutic efficacy of PD-1/PD-L1 inhibitors (22,44,45). The level of PD-L1 expression can be assessed by tumor proportion score (TPS), which is defined as the tumor percentage of viable tumor cells showing partial or complete membrane staining and CPS. It is evaluated based on the total number of PD-L1 positive cells (tumor, lymphocytes, and macrophages) compared to total tumor cells; hence, the factors from both tumor and immune cells can be considered comprehensively. Several studies have suggested that patients with tumors of high CPS or TPS benefit from pembrolizumab treatment. In the KEYNOTE-180 trial, positive PD-L1 expression (defined as CPS ≥10) in esophageal tumors was 47.9% (58/121). Patients with PD-L1$^+$ expression had a higher 6-month PFS rate (22% vs. 10%) and 9-month PFS rate (14% vs. 5%) than those with PD-L1$^-$ expression (22). In a phase III KEYNOTE-181 study of pembrolizumab vs. the investigator’s choice of chemotherapy as second-line therapy for patients with advanced/metastatic ESCC, EAC, or esophagogastric Junction (EGJ) (NCT02564263), pembrolizumab was superior to chemotherapy in terms of OS in patients with PD-L1 CPS ≥10 (Hazard ratio 0.69; 95% CI: 0.52–0.93; P=0.0074). The 12-month OS rate in patients with CPS ≥10 was 43% vs. 20% (23).

Some studies have also indicated a poor association between PD-L1 and clinical response. Huang et al. found that in 24 patients with metastatic ESCC, 15 (62.5%) had PD-L1$^+$ tumors (defined as ≥5%). The patients with PD-L1$^+$ tumors tended to have higher ORR (20%, 2/10) and PD-L1$^-$ patients (25%, 6/24) (29).
Of the various anti-PD-L1 antibodies available for testing PD-L1 expression, four antibodies (22C3 and 28-8 from Dako; SP263 and SP142 from Ventana) have been used widely in clinical trials. Despite these antibodies recognizing different peptide-recognizing domains in PD-L1 antigen, their testing efficacy has been shown to be consistent apart from a relatively lower positive detection rate in SP142. Because of antigen degradation during preservation and heterogeneous expression in tumor tissues across different locations, the preservation time and specimen size (surgery vs. biopsy) should also be considered when PD-L1 expression is evaluated.

**TMB**

There is a significant correlation between the TMB and the response to PD-1/PD-L1 inhibitors in a wide variety of cancers (17). EC has an upper middle level of TMB in solid tumors (46,47). Huang et al. performed exome sequencing on the DNAs from paired tumors and white blood cells from 23 patients with metastatic ESCC (28). The mutation count per tumor ranged from 15 to 219, with a median of 60 mutations per tumor. Using 60 mutations per tumor as the cut-off value, 8 of 11 samples with high TMB showed clinical benefit and only 2 of the 12 samples with low TMB had clinical benefit (P=0.0123). Further explorations of TMB in EC treated with immunotherapy are ongoing. Although TMB is an important feature correlated with immunotherapy, it is time-consuming and requires a higher quality of specimens compared with IHC-based PD-L1 assessment. TMB will not replace PD-L1 test, but could be a beneficial supplement.

**MSI**

DNA replication errors are checked and corrected by MMR system. MMR proteins including MLH1, PMS2, MSH2, and MSH6 can form heterodimers to correct such mismatches. Deficient mismatch repair (dMMR) in tumor cells causes areas of DNA to become unstable, resulting in MSI. Tumors with a high level of MSI (MSI-H) are likely to have a high TMB count and an increased number of tumor neoantigens. Thus, powerful local and systemic immune responses might be induced (48). Chalmers et al. analyzed MSI-H and TMB in 62,150 patients with solid tumors and found that the vast majority of MSI-H samples also had high TMB (83%) and 97% had TMB ≥10 mutations/Mb (46). A clinical trial of PD-1 blockade in patients with advanced MMR-deficient in 12 different types of tumor showed that 53% and 21% of patients achieved objective radiographic response or complete response, respectively (49). Other results from 5 clinical trials including 15 tumor entities with MSI/dMMR tumors (KEYNOTE-012, -016, -028, -158 and -164) also revealed a high correlation between MSI and a better response to ICIs. These findings led the FDA to approve a tumor-type agnostic pembrolizumab for all

| Table 3 A list of biomarkers in immune checkpoint inhibitor therapy for ESCC |
|---|
| Factor | Association with favorable clinical outcome | Tissue type for biomarker assessment | Possible assay type for biomarker assessment | Cutoff value (%) | Positive rate (%) |
|---|---|---|---|---|---|
| PD-L1 expression | Positive | Tissue | IHC | 1% or 10% for CPS | 41.4% |
| Tumor mutational burden | High | Tissue/blood (ctDNA) | WES or targeted sequence | 7.7 | Media |
| MSI-H or dMMR | MSI-H or dMMR | Tissue | IHC or WES | NA | 4.35% |
| Copy number variation | High/low | Tissue | WES or targeted sequence | NA | NA |
| T cell repertoire clonality change | Clonality/diversity | Tissue/blood | Multiple PCR or sequence | NA | NA |
| T cell-inflamed microenvironment | High | Tissue | PCR | Hot tumor | 11.7% |

ESCC, esophageal squamous cell carcinoma; IHC, immunohistochemistry; CPS, combined positive score; MSI-H, high microsatellite instability; dMMR, deficient mismatch repair; PCR, pathologic complete response.
microsatellite-unstable tumors.

The dMMR in esophageal carcinoma stands at 4.35% (47). In the KEYNOTE-012 study, 24 samples of gastric or EGJ cancer were analyzed, and 17% of which were MSI-H tumors. In the CHECKMATE-032 study, in which patients with esophageal, EGJ, or gastric cancer were included, MSI-H occurred in 18% of samples (26). MSI is rarely reported in ESCC and is a potential area of focus for future studies.

**T-cell-inflamed gene-expression profile (GEP)**

The ability of the host immune system to recognize and eliminate tumor cells is orchestrated by innate and adaptive antitumor immune response. IFN-γ is the key cytokine and plays a vital role in the antitumor response. Meanwhile, as a feedback mechanism, IFN-γ signaling activates the PD-1 signaling axis via the upregulation of PD-L1 and PD-L2 in tumor cells as well as in stromal cells. Multigene immune signatures might comprehensively capture a complex, T cell-inflamed phenotype necessary for the clinical activity of PD-1/PD-L1 inhibitors.

In a pilot study involving 19 melanoma patients and, eventually, 220 patients with 9 cancers from KEYNOTE-012 and -028, a T cell-inflamed GEP comprising 18 genes was found to be associated with response to pembrolizumab, independently based on PD-L1 expression (50). The GFP genes selected were related to antigen presentation, chemokine expression, cytotoxic activity, and adaptive immune resistance, and were identical to the 18 genes in the tumor inflammation signature (50,51). Later, the KEYNOTE-028 study of 313 patients consistently found higher T-cell-inflamed GFP score in patients who achieved improved ORR and longer PFS after pembrolizumab treatment (44). In addition to these two studies, an analysis of 23 patients with EC (including 8 patients with ESCC) from KEYNOTE-028 used an IFN-γ GFP signature that included 6 genes of the 18-gene profile. The result also suggested that the patients with higher IFN-γ composite scores achieved better treatment outcomes (21).

**TILs**

During tumor progression and the application of various therapies, lymphocytes migrate to the tumor tissues to serve as effectors of antitumor response. At the same time, because of the inhibitory tumor microenvironment or feedback mechanism, TILs are usually inactivated. Since ICIs target inhibitory checkpoint molecules, antitumor response might be reactivated in tumors with high levels of TILs. An early study of metastatic melanoma suggested that TILs were predictive of response in both CTLA-4 and PD-1 blockade. TILs were found at a higher density early during treatment in patients who showed a better response (52). A study examined the prognostic impact of PD-L1 expression and TIL status, and their combination, in EC. Of the 305 patients included, 279 (91%) had ESCC. The results indicated that both PD-L1 positivity and TIL positivity were significantly correlated with better OS and DFS. There was also a close association observed between TIL positivity and high expression of CD8. Interestingly, the combination of PD-L1 expression and TIL status could further classify the patients according to clinical outcome: the patients in the PD-L1 TIL+ group had the longest OS and DFS, and those in the PD-L1 TIL− group had the shortest OS and DFS (53). Patients with PD-L1 TIL+ tumors were the most suitable candidates to receive mono-PD-1/PD-L1 inhibitors. Patients with PD-L1 TIL− tumors would not benefit from PD-1/PD-L1 blockade. For patients with either PD-L1 TIL− or PD-L1 TIL+ tumors, strategies to attract TILs are needed first before any ICI treatment is performed (53).

**Other biomarkers**

A recent study in advanced melanoma indicated that the functional immunological assay on ex vivo tumor-metastatic lymph nodes could serve as a new predictor in immunotherapy (54). Circulating immune cells, which are more convenient and can be obtained less invasively than tumor tissue, are potential biomarkers for predicting treatment response to ICIs. PD-L1 expression on peripheral T cells was associated with the response to anti-CTLA-4 mAb (54). Meanwhile, CD137 expression on circulating CD8+ T cells was discovered to be a promising predictor after adjuvant ipilimumab combined with nivolumab (54). Circulating tumor DNA (ctDNA) after CRT has been shown to be associated with tumor progression, metastasis, and disease-specific survival in patients with EC (55). Improved antitumor immunity and response were found in melanoma patients with favorable gut microbiomes (55). Moreover, another study showed that gut microbiome promoted the efficacy of ICIs through the recruitment
of T cells into the tumor beds (56). Although the role of esophageal microbiomes in ESCC patients is not well known, a study including 25 ESCC patients and 50 matched controls found that Prevotella, especially Prevotella nanceiensis, was abundant in patients with ESCC (57). Furthermore, another case-control study on 210 esophageal samples from 86 patients suggested that there was decreased microbial diversity in EAC tissues compared with tissues from healthy control patients (58). The association between the esophageal microbiome and the treatment response to ICIs requires further study in future investigations.

Because of intratumoral heterogeneity, the levels of biomarkers from biopsies obtained from different sites might be inconsistent in a single patient. The fluctuating levels of biomarkers may also be attributed to individual genetic background. Different therapeutic strategies and alterations of the tumor microenvironment and systemic immunity during treatment result in dynamic changes in some predictive biomarkers, such as PD-L1 expression, TILs, and peripheral immune cells. Furthermore, a lack of consistency between different biomarkers might be explained by the different mechanisms involved (59). In TMB, specific mutations in oncogenes and tumor PD-L1 expression describe the tumor component while immune-cell PD-1/PD-L1 expression, HLA genotype, TCR repertoire, and possibly immune signatures represent the antitumor immune response. Therefore, a combination of several biomarkers with different mechanism could achieve enhanced ability to predict the clinical response and treatment outcome.

Conclusions and future directions

The findings of clinical trials have emphasized that the combination of ICIs with chemotherapy or RT might achieve greater treatment efficacy for various cancers compared to the use of ICIs alone. The studies of anti-PD-1/L1 antibody combined with CRT in ESCC are ongoing. The exploration of strong predictive and prognostic biomarkers or integrated biomarkers to optimize therapeutic strategies is another area of focus for research. Meanwhile, the identification of valuable biomarkers for predicting toxicity is also crucial. With a deepening understanding of the biological tumor cells and antitumor immune features in ESCC, it is reasonable to believe that ICIs, or their application in combined treatment, will benefit more patients in future.

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