Emerging PD-1/PD-L1 targeting immunotherapy in non-small cell lung cancer: Current status and future perspective in Japan, US, EU, and China

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Non-small cell lung cancer (NSCLC), one of the deadliest types of cancers worldwide, has been the target of immunotherapy due to its high immune antigenicity. With the addition of immune-checkpoint inhibitors (ICIs), including anti-PD-1/PD-L1 antibodies, as an indispensable and powerful regimen for the treatment of this lethal disease, the median survival time for patients with stage IV NSCLC is approximately 2 years. In contrast, the response rate to ICIs remains less than 50%, even if the patients are selected using biomarkers such as PD-L1. Pharmaceutical companies have begun to develop additional anti-PD-1/PD-L1 antibodies to overcome resistance and are devising further immunotherapy combinations. More than 20 anti-PD-1/PD-L1 antibodies have been approved or are currently in development. Numerous combination therapies are under development, and several combination therapies have provided positive results in randomized controlled trials. This review aimed to examine the current status of approved and investigational anti-PD-1/PD-L1 antibodies for NSCLC in Japan, the United States, the European Union, and China. Further, this review discusses the challenges and future perspectives for developing new ICIs in alignment with the global developments in Japan.

KEYWORDS
non-small cell lung cancer, immunotherapy, anti-PD-1 antibody, anti-PD-L1 antibody, clinical trial
1 Introduction

Nivolumab was approved as a treatment for malignant melanoma by the United States (US) Food and Drug Administration (FDA) in 2014. With this approval, immune-checkpoint inhibitors (ICIs), mainly anti-programmed cell death-1 (PD-1) antibodies, have rapidly revolutionized cancer treatment. ICIs and anti-PD-1 antibodies have become the main component of cancer therapy and are used in all types of cancer (1). In Japan, since the approval of the first anti-PD-1 antibody nivolumab in 2015, pembrolizumab, the anti-PD ligand (PD-L)1 antibodies atezolizumab, durvalumab, and avelumab; and the anti-cytotoxic T-lymphocyte-associated protein 4 (CTLA-4) antibody ipilimumab have also been approved and reimbursed by the National Health Insurance system. Among all the drugs available in Japan in terms of sales as of 2021, pembrolizumab and nivolumab rank first and second, respectively, with each drug generating more than $1 billion in sales (2).

In non-small cell lung cancer (NSCLC), according to the NCCN Guidelines (version 1.2022) (3) and the Japanese Lung Cancer Society Guidelines for NSCLC (2021 edition) (4), combination therapy with an anti-PD-1/PD-L1 antibody and platinum-based chemotherapy is recommended as first-line therapy for patients with advanced NSCLC without driver gene mutations. Even if an anti-PD-1/PD-L1 antibody is not used in the initial therapy, it is recommended as a second-line or subsequent therapy. In locally advanced lung cancer, durvalumab is recommended as maintenance therapy after chemoradiation therapy, based on the results of the PACIFIC trial (5). In terms of postoperative adjuvant chemotherapy, sequential atezolizumab after adjuvant chemotherapy was approved on October 2021 by the FDA and recommended in the NCCN guidelines as the first perioperative ICI for PD-L1-positive (≥1%) stage II-IIIA NSCLC after radical resection based on the results of the IMpower010 trial (6). In Japan, an application for partial changes to atezolizumab as adjuvant therapy was submitted in July 2021 and approved in May 2022. The ICIs must be considered in all patients with NSCLC, except for those with a positive driver gene mutation, poor performance status, autoimmune disease, or interstitial lung disease.

Currently, ICIs primarily target PD-1/PD-L1, which are immune-checkpoint molecules that inhibit the priming phase (activation of antigen-presenting cells and T cells) and effector phase (direct damage to cancer cells) of the cancer-immunity cycle (7). In 2011, 3 years prior to the approval of nivolumab, an antibody drug targeting CTLA-4 (ipilimumab; another immune-checkpoint molecule primarily involved in the priming phase), was developed and approved by the FDA for the treatment of unresectable or metastatic malignant melanoma (approved in Japan in 2015) (8, 9) (Figure 1). Ipilimumab is currently approved in the United States (US), European Union (US), and Japan in combination with nivolumab (with or without platinum combination chemotherapy) as first-line treatment for NSCLC, regardless of the PD-L1 expression status (7, 13, 14). The Japan Clinical Oncology Group is currently conducting a phase III study (JCOG2007, NIPPON study) to evaluate the superiority of nivolumab plus ipilimumab plus platinum combination chemotherapy over pembrolizumab plus platinum combination chemotherapy, which has attracted research attention as an optimal first-line treatment for patients with NSCLC (15).

Currently, multiple ICIs are approved, and other pharmaceutical companies have developed their own anti-PD-1/PD-L1 antibodies as monotherapy and combination immunotherapy for new indications. However, no study has provided a summary of the entire picture of the early and late phase of ICI development from a global perspective. This review aimed to provide a comprehensive overview and a better understanding of the emerging anti-PD-1/PD-L1 antibodies for NSCLC. We also aimed to discuss the current challenges and future perspectives on the development of ICIs in Japan.

2 Immunotherapies in development

2.1 Novel PD-1/PD-L1 pathway inhibitors

The anti-PD-1 antibodies nivolumab and pembrolizumab and the anti-PD-L1 antibodies atezolizumab, durvalumab, and avelumab (not indicated for NSCLC) are available worldwide, and pharmaceutical companies are developing more anti-PD-1/PD-L1 antibodies with the aim of obtaining additional indications for diseases for which anti-PD-(L)1 antibodies are not available or as combined immunotherapies with compounds that they are developing. The details of the anti-PD-1/PD-L1 antibodies currently under development are presented in Table 1.

2.1.1 Cemiplimab

Cemiplimab is an anti-PD-1 antibody approved by the FDA in September 2018 for the treatment of cutaneous squamous cell carcinoma (cSCC), and in February 2021 for cutaneous basal cell carcinoma (cBCC) and for first-line treatment of NSCLC with PD-L1 expression of ≥50% (47, 48). The approval of its use was based on the results of the EMPOWER-Lung 1 study, a phase III trial that compared the efficacy of cemiplimab alone with that of platinum-based chemotherapy in patients with advanced NSCLC with tumor PD-L1 (22C-3) expression of ≥50% with overall survival (OS) and progression-free survival (PFS) as the primary endpoints. The median OS times of the cemiplimab and chemotherapy group were 22.1 months (95% confidence interval [CI], 17.7–21.4) and 14.3 months (95% CI, 11.7–19.2), respectively, with a hazard ratio (HR) of 0.68 (95% CI, 0.53–
TABLE 1 Anti-PD-1/PD-L1 antibody monotherapy currently approved and under development (data on March 31, 2022).

| Drug name | Target | FDA | EMA | PMDA | NMPA | Approval status in NSCLC | Other major approved indications | Pivotal trial | R&D institutions | Country | Business partner | Reference |
|-----------|--------|-----|-----|------|------|--------------------------|---------------------------------|--------------|------------------|---------|-----------------|----------|
| Pembrolizumab | PD-1 | Q1L (TPS ≥ 1%), 2L | Melanoma, HNSCC, chL, urothelial cancer, gastric cancer, cervical cancer, PMBC, HCC, MCC, esophageal cancer, cSCC, MSI-H/MMR/ TMB-H solid tumor, RCC* | KEYNOTE-024, KEYNOTE-042, KEYNOTE-010 | Merck | Germany | (16–19) |
| Nivolumab | PD-1 | Q2L | Melanoma*, RCC, chL, gastric cancer, HNSCC, urothelial cancer, MSI-H/MMR CRC, HCC, esophageal cancer* | CheckMate-017, CheckMate-057 | Ono Pharmaceutical | Japan | Bristol-Myers Squibb | (20–22) |
| Atezolizumab | PD-1 | Q1L (TCJ or IC3), 2L (adjunct use is approved by the FDA and the PMDA) | Urothelial cancer | Ipilimumab, Nivolumab | Boehringer Ingelheim | Switzerland | Chugai Pharmaceutical | (6, 23, 24) |
| Durvalumab | PD-1 | Q stage III after CRT | SCLC | PACIFIC | MedImmune | US | AstraZeneca | (5, 25) |
| pembrolizumab | PD-1 | Q1L (TPS ≥ 50%) | cSCC, basal cell carcinoma | EMPOWER-2 | Regeneron | US | Sanofi | (26, 27) |
| sintilimab | PD-1 | × | × | Q≤5q 2L | chL, HCC | ORIENT-3 | Innovon Biologies | China | Eli Lilly | (28) |
| Tislelizumab | PD-1 | × | × | Q2L | cHIL, HCC, esophageal cancer, nasopharyngeal cancer, urothelial cancer, MSI-H/MMR solid tumor | RATIONALE 303 | BeiGene | China | Novartis | (29) |
| Sugemalimab | PD-L1 | × | × | × | Q stage III after CRT | GEMSTONE 301 | cStonest Pharmaceuticals | China | Pfizer | (30, 31) |
| Camrelizumab | PD-1 | × | × | × | chL, HCC, esophageal cancer | Jiangsu Hengrui Pharmaceuticals | China | LSK BioPharma | (32) |

(Continued)
while the median PFS times were 8.2 months (95% CI, 6.1–8.8) and 5.7 months (95% CI, 4.5–6.2) months, respectively, with an HR of 0.54 (95% CI, 0.43–0.68; p < 0.0001), leading to the early discontinuation of the trial due to its significant superiority (26). The EMPOWER-Lung 3 trial (NCT03409614), a phase III trial that evaluated the superiority of cemiplimab over platinum-based chemotherapy, was also discontinued early as it already showed substantial evidence of cemiplimab’s significant superiority over other therapy; hence, an application for the approval of its use as first-line treatment for NSCLC has been submitted to the FDA (49). However, pivotal studies of cemiplimab for cBCC, cSCC, and NSCLC did not include Japanese patients; hence, cemiplimab has not yet been approved in Japan. Several studies on cemiplimab are underway in Japan (NCT03257267 and NCT03969004), and the results of these trials are expected to contribute to its approval.

### 2.1.2 Sintilimab

In China, which has been a member of the International Council for Harmonization of Technical Requirements for Pharmaceuticals for Human Use since 2017 and became a member of the Management Board in 2018, several anti-PD-1/PD-L1 antibodies developed by domestic companies have been tested in clinical trials and approved for reimbursement. Anti-PD-1 antibodies, such as sintilimab, tislelizumab, toripalimab, zimberelimab, penpulimab, and camrelizumab, and anti-PD-1 antibodies, such as sugemalimab and socazolimab, have been approved in China at relatively lower prices compared with those developed by pharmaceutical companies outside China, and are cost-effective compared with chemotherapies (50–53). In recent years, several Chinese companies have collaborated with overseas companies to expand their businesses worldwide.

In the ORIENT-11 study, a phase III trial that evaluated the superiority of sintilimab plus platinum-based chemotherapy...
over platinum-based chemotherapy as first-line treatment for advanced non-squamous NSCLC, the PFS times were 8.9 months (95% CI, 7.1–11.3) in the sintilimab group and 5.0 months (95% CI, 4.8–6.2) in the platinum combination chemotherapy group, with an HR of 0.482 (95% CI, 0.36–0.64; p < 0.00001), showing the superior efficacy of sintilimab combination therapy over platinum-doublet therapy (54). Based on the results of a biomarker analysis reported at the 2021 World Conference on Lung Cancer, the abundant expression of genes in the major histocompatibility complex class II antigen-presenting pathway was predictive of the response to sintilimab plus chemotherapy, regardless of the PD-L1 expression status, and is a predictor of treatment response, in addition to tumor PD-L1 expression and tumor mutational burden (55). For patients with squamous NSCLC, the Phase III ORIENT-12 trial compared the efficacy of sintilimab plus gemcitabine plus platinum-based chemotherapy with that of platinum-doublet therapy as first-line treatment, with PFS times of 5.5 months and 4.9 months and an HR of 0.536 (p < 0.00001), showing the significant superiority of the sintilimab combination (36). Based on the results of these trials, Eli Lilly and Sintilimab’s development partners were in the process of submitting an application to the FDA for the approval of its use. However, on February 2022, the FDA dismissed this request and required additional clinical studies because these trials used PFS rather than OS as their primary endpoint; trials conducted only in China did not reflect the racially diverse population of the US, and the FDA was previously consulted regarding the study design, endpoints, and control group selection (57).

2.1.3 Tislelizumab

Tislelizumab, another anti-PD-L1 antibody that has shown efficacy in phase III trials, was designed to minimize macrophage binding to the Fc gamma receptor. This mechanism is believed to suppress the antibody-dependent cell-mediated phagocytosis, which can lead to T-cell elimination and resistance to PD-L1 inhibitors (58–60). The RATIONALE 304 study, an open-label phase III trial conducted in China, evaluated the superiority of tislelizumab plus pemetrexed plus platinum-based chemotherapy over standard platinum-doublet therapy as first-line treatment of non-squamous NSCLC, which reported significantly better median PFS in the tislelizumab group (9.7 months) compared with that in the control arm (7.6 months), with an HR of 0.645 (p = 0.0044) (61). In the RATIONALE 307 study, a three-arm, open-label phase III trial conducted in China for the first-line treatment of squamous NSCLC, the experimental arms received paclitaxel plus carboplatin or nab-paclitaxel plus carboplatin combined with tislelizumab, while the control arm received chemotherapy with paclitaxel plus carboplatin or nab-paclitaxel plus carboplatin. The median PFS times were 7.6 months, 7.6 months, and 5.5 months, respectively, with a significantly better HR in the tislelizumab combination group compared with that in the control arm, indicating that tislelizumab is effective regardless of the PD-L1 expression status (62). A biologics license application for tislelizumab was submitted to the FDA on September 2021 by Novartis, which has development and commercialization rights in North America, Europe, and Japan, based on the results of the RATIONALE 302 study, a global phase III trial on esophageal squamous cell cancer (63). However, the tislelizumab trial on NSCLC was conducted only in China, and it may be difficult to obtain an FDA approval based solely on the current results, similar to the case of sintilimab. A global phase III study (NCT04746924) evaluating the superiority of tislelizumab in combination with ociperlimab (an anti-TIGIT antibody) over pembrolizumab in advanced NSCLC patients with PD-L1 expression of ≥50% and a phase III study (NCT04866601) evaluating the superiority of tislelizumab in combination with ociperlimab over durvalumab after chemoradiotherapy are underway, and further developments are anticipated.

2.1.4 Subcutaneous PD-1/PD-L1 inhibitors

As some patients with cancer who received ICIs achieve long-term survival and are able to continue receiving them for extended periods of time, companies are also developing ICIs that can be administered subcutaneously in the pursuit of improving convenience during long-term administration and to gain approval for alternative routes of administration. Not only the subcutaneous formulations of approved ICIs, such as nivolumab (NCT03656718), pembrolizumab (NCT04956692), atezolizumab (NCT03735121), and durvalumab (NCT04870112), but also newer agents such as sasanlimab and envafolimab have completed phase I trials and are in further development without any notable safety differences compared with the intravenous formulations, and it will not take long before they are approved and become part of our treatment options (43, 45, 64).

2.2 Novel combination therapies with PD-1/PD-L1 pathway inhibitors

Cancer immunotherapies with monoclonal antibodies that inhibit the PD-1 pathway have significantly impacted the treatment of patients with cancer in recent years. However, despite the remarkable clinical efficacy of these agents, anti-PD-L1 monotherapies are not actively used in many patients. Evidence from the combined inhibition of PD-1 and CTLA-4 in melanoma and NSCLC underscores the potential of combining drugs with synergistic mechanisms of action to further enhance the clinical efficacy of monotherapies, which has encouraged the ongoing development of combination therapies with ICIs, including anti-CTLA-4 antibodies and
chemotherapy (65). The currently approved and investigated anti-PD(L)1 combination therapies are listed in Table 2.

2.2.1 CTLA-4 inhibitors

The most advanced combination therapy with anti-PD-(L)1 antibodies is ipilimumab, an anti-CTLA-4 antibody developed by Bristol-Myers Squibb. Ipilimumab is currently approved in Japan for use in combination with nivolumab for the treatment of NSCLC, malignant pleural mesothelioma, malignant melanoma, renal cell carcinoma, and colorectal cancer with high-frequency microsatellite instability. It is the only anti-CTLA-4 antibody approved for NSCLC patients in Japan and other countries based on the results of CheckMate-227 and -9LA studies (69, 70). The CheckMate-227 randomized, open-label, phase 3 trial, enrolled patients with stage IV or recurrent NSCLC. Patients with a PD-L1 expression level of 1% were randomly assigned in a ratio of 1:1:1 to receive nivolumab plus ipilimumab, nivolumab alone, or chemotherapy. Patients with a PD-L1 expression level of less than 1% were randomly assigned in a 1:1:1 ratio to receive nivolumab plus ipilimumab, nivolumab plus chemotherapy, or chemotherapy alone. The median OS times in the nivolumab plus ipilimumab combination group were significantly superior to those in the chemotherapy group, regardless of PD-L1 expression level. In the CheckMate-9LA randomized, open-label, phase 3 trial, patients with treatment-naïve advanced NSCLC were assigned to receive nivolumab (360 mg intravenously every 3 weeks) plus ipilimumab (1 mg/kg intravenously every 6 weeks) combined with histology-based, platinum-doublet chemotherapy (intravenously every 3 weeks for two cycles; combination group), or chemotherapy alone (every 3 weeks for four cycles; control group). The median OS was 15.6 months (95% CI, 13.9-20.0) in the experimental group versus 10.9 months (95% CI, 9.5-12.6) in the control group (HR=0.66 [95% CI, 0.55-0.80]). In contrast, tremelimumab, an anti-CTLA-4 antibody, has been granted an Orphan Drug Designation for malignant pleural mesothelioma by the FDA in 2015 and has been primarily used in combination with durvalumab as treatment for advanced cases. However, in the

TABLE 2 Anti-PD-1/PD-L1 antibody combination therapy currently approved and under development (data on March 31, 2022).

| Drug combination | Target | FDA | EMA | PMDA | NMPA | Other major approved indications | Pivotal trial | R&D institutions | Country | Business partner | Reference |
|------------------|--------|-----|-----|------|------|----------------------------------|--------------|------------------|---------|------------------|----------|
| Pembrolizumab + chemotherapy | PD-1 | ○ | ○ | ○ | ○ | RCC, endometrial cancer, esophageal cancer, gastric cancer, cervical cancer, breast cancer* | KEYNOTE-189, KEYNOTE-407 | Merck | Germany | (19, 66, 67) |
| Atezolizumab + chemotherapy | PD-L1 | ○ | ○ | ○ | ○ | SCLC, HCC, melanoma | IMpower-130 | Roche | Switzerland | (24, 68) |
| Nivolumab + ipilimumab | PD-1 | ○ | ○ | ○ | ○ | Melanoma, RCC, MSI-H/DMMR CRC, HCC, malignant mesothelioma | CheckMate-227 | Ono Pharmaceutical | Japan | Bristol-Myers Squibb | (22, 69) |
| Nivolumab + ipilimumab + chemotherapy | PD-1 | ○ | ○ | ○ | × | – | CheckMate-9LA | Ono Pharmaceutical | Japan | Bristol-Myers Squibb | (22, 70) |
| Nivolumab + chemotherapy | PD-1 | ○* | × | × | × | Gastric cancer | CheckMate-816 | Ono Pharmaceutical | Japan | Bristol-Myers Squibb | (71) |
| Complimab + chemotherapy | PD-1 | × | submitted | × | × | – | EMPower-Lung 3 | Regeneron | US | Samson | (49) |
| Sintilimab + chemotherapy | PD-1 | × | × | × | ○ | HCC | ORIENT-11, ORIENT-12 | Innovent Biologies | China | Eli Lilly | (35, 56) |
| Camrelizumab + chemotherapy | PD-1 | × | × | × | ○ | Nasopharyngeal cancer | Camel, Camel-Sq | Jiangsu Hengrui Pharmaceuticals | China | (72, 73) |
| Tislelizumab + chemotherapy | PD-1 | × | × | × | ○ | HCC, esophageal cancer, nasopharyngeal cancer | RATIONALE 304, RATIONALE 307 | BeGene | China | Novartis | (61, 62) |
| Sugemalimab + chemotherapy | PD-L1 | × | × | × | ○ | – | GEMSTONE 302 | Genetech Pharmaceuticals | China | Pfizer | (74) |
| Toripalimab + chemotherapy | PD-1 | × | × | × | × | Esophageal cancer, nasopharyngeal cancer | CHOICE-01 | Shanghai Junshi Biosciences | China | Coherus Biosciences | (75) |

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TABLE 2 Continued

| Drug combination | Target | FDA | EMA | PMDA | NMPA | Other major approved indications | Pivotal trial | R&D institutions | Country | Business partner | Reference |
|------------------|--------|-----|-----|------|------|---------------------------------|--------------|-----------------|---------|-----------------|-----------|
| Avelumab + chemotherapy | PD-L1 | × | × | × | × | RCC | Merck | Germany | Pfizer | (76, 77) |
| Pembrolizumab + chemotherapy | PD-1 | × | × | × | × | | Akerso | China | | (78) |
| Retifanlimab + chemotherapy | PD-1 | × | × | × | × | | MacroGenics | US | Incyte | (79) |
| Serplulimab + chemotherapy | PD-1 | × | × | × | × | | Shanghai Henlius Biotech | China | PT Kalbe Genexine Biologics | (80, 81) |
| Cosbelimab + chemotherapy | PD-L1 | × | × | × | × | | Checkpoint Therapeutics | US | | (82) |
| Ezalimumab + investigational drugs | PD-1 | × | × | × | × | | Boehringer Ingelheim | Germany | | (83–85) |
| Spartalizumab + investigational drugs | PD-1 | × | × | × | × | | Novartis | Switzerland | | (86) |
| Geptanolimab + fruquintinib | PD-1 | × | × | × | × | | Genor Biopharma | China | Apollomics | (87) |
| Sasnulimab + investigational drugs (subcutaneous) | PD-1 | × | × | × | × | | Pfizer | US | | (88) |
| Pembrolizumab + chemotherapy (subcutaneous) | PD-1 | × | × | × | × | | Merck | Germany | | (89) |

CRC, colorectal cancer; cSCC, cutaneous squamous cell carcinoma; dMMR, deficient mismatch repair; EMA, European Medicines Agency; FDA, Food and Drug Administration; HCC, hepatocellular carcinoma; MSI-H, microsatellite instability-high; NMPA, National Medical Products Administration; PMBC, primary mediastinal B-cell lymphoma; PMDA, Pharmaceuticals and Medical Devices Agency; RCC, renal cell carcinoma; SCLC, small-cell lung cancer; TMB-H, tumor mutational burden-high.

*Approval as neoadjuvant therapy.

MYSTIC study, a phase III trial on NSCLC, the efficacy of combination of durvalumab and tremelimumab was compared with that of platinum doublet chemotherapy but failed to achieve the co-primary endpoints, with median OS times of 11.9 months (95% CI, 9.0–17.7) and 12.9 months (95% CI, 10.5–15.0), respectively, with an HR of 0.85 (95% CI, 0.61–1.17; p = 0.20), and median PFS times of 3.9 months (95% CI, 2.8–5.0) and 5.4 months (95% CI, 4.6–5.8), respectively, with an HR of 1.05 (95% CI, 0.72–1.53; p = 0.71); it has not yet been approved for use in Japan or any other country (90). The results of a phase III trial (POSEIDON study) that evaluated the superiority of tremelimumab in combination with durvalumab and platinum-based chemotherapy over platinum-based chemotherapy as first-line treatment for NSCLC was presented at the 2021 World Conference on Lung Cancer, with favorable outcomes: median OS times of 14.0 months (95% CI, 11.7–16.1) and 11.7 months (95% CI, 10.5–13.1), respectively, with an HR of 0.77 (95% CI 0.63–0.92; p = 0.00304), and median PFS times of 6.2 months (95% CI, 5.0–6.5) and 4.8 months (95% CI, 4.6–5.8), respectively, with an HR of 0.72 (95% CI, 0.60–0.86; p = 0.00031) (91). The primary endpoints tremelimumab in combination with durvalumab and chemotherapy were PFS and OS. The OS was not met, and the efficacy of durvalumab in combination with tremelimumab and chemotherapy was analyzed as the pre-specified key secondary endpoint. In its press release, AstraZeneca stated, “We look forward to discussing these data with regulatory authorities.” However, whether an application was actually filed remains unclear (92). Other drugs targeting CTLA-4, including an open-label phase II study of NSCLC patients treated with KN046, a recombinant humanized PD-L1/CTLA-4 bispecific antibody, were reported during the 2021 meeting of the American Society of Clinical Oncology (ASCO). Based on the hypothesis that the limited peripheral distribution of KN046 would reduce the incidence of treatment-related toxicity, KN046 was added to platinum combination chemotherapy as a treatment for squamous and non-squamous NSCLC and showed good safety and promising efficacy (93). KN046 is currently under phase III trials in combination with carboplatin plus paclitaxel (NCT04474119) as a first-line treatment for squamous NSCLC and in combination with lenvatinib (NCT05001724) for NSCLC after ICI resistance, and is expected to show efficacy and safety as a novel CTLA-4 inhibitor.
2.2.2 LAG-3 inhibitors

In 2021, a phase III study (RELATIVITY-047 trial) comparing the efficacy of nivolumab combined with the anti-lymphocyte-activation gene 3 (LAG-3) antibody relatlimab with that of nivolumab alone as first-line treatment for malignant melanoma was the first to show the significant benefit of adding an anti-LAG-3 antibody to standard immunotherapy. The median PFS times, the primary endpoint of the study, were 10.1 months (95% CI, 6.4–15.7) and 4.6 months (95% CI, 3.4–5.6), respectively, with an HR of 0.75 (95% CI, 0.60–0.90; p = 0.0055) (94). The results were presented at ASCO 2021, and relatlimab was approved by the FDA as treatment for unresectable or metastatic melanoma on March 18, 2022, and was included in the PD-1/PD-L1 and anti-CTLA-4 antibody combination with pembrolizumab (NCT03516981). LAG-3 is a cell surface molecule that is expressed in effector and regulatory T cells and regulates the T-cell response, activation, and proliferation. Inhibition of the LAG-3 pathway restores the exhausted T-cell function and promotes antitumor responses, and the use of LAG-3 with PD-1/PD-L1 pathway inhibitors as combination therapy is expected. In NSCLC, several potential novel drug combinations are currently under investigation, including phase II trials of platinum-based chemotherapy plus nivolumab in combination with relatlimab as first-line therapy (NCT04623775); efalagimod alpha, a soluble fusion protein of LAG-3 and the human IgG Fc moiety, in combination with pembrolizumab as treatment for patients who showed resistance to anti-PD-1/PD-L1 therapy (NCT03625323); and favezelimab, an anti-LAG-3 antibody, in combination with pembrolizumab (NCT03516981).

2.2.3 TIGIT inhibitors

Along with LAG-3, T-cell immunoreceptors with immunoglobulin and ITIM domains (TIGIT) are new candidate ICIs. TIGIT suppresses T-cell activation, exhausts T cells, and is highly expressed in tumor-infiltrating T cells. The inhibition of TIGIT promotes cytotoxic T-cell proliferation and antitumor responses, leading to the development of combination therapy with anti-PD-1/PD-L1 antibodies (96, 97). The FDA granted the breakthrough therapy designation to tiragolumab, an anti-TIGIT antibody, in combination with atezolizumab for the treatment of NSCLC with high PD-L1 expression on January 2021 according to the results of the CITYSCAPE trial, a randomized, double-blind, placebo-controlled phase II trial that examined the efficacy and safety of tiragolumab in combination with atezolizumab as first-line therapy in NSCLC patients with PD-L1 expression of ≥1%, which was presented at ASCO 2020 (98). Tiragolumab plus atezolizumab was able to achieve the co-primary endpoints in the intention-to-treat population, showing an improvement in the overall response rate (ORR) (37% vs. 21%) and PFS (median PFS, 5.6 vs. 3.9 months; HR, 0.58; 95% CI, 0.38–0.89) compared with atezolizumab alone. In the PD-L1 expression of ≥50% sub-population, the ORRs were 66% and 24%, while the PFS times were not reached and 4.1 months, respectively, with an HR of 0.30 (95% CI, 0.15–0.61), showing very favorable results in the combination therapy group with an acceptable safety profile. Based on these results, Roche conducted a phase III trial (SKYSCRAPER-01, NCT04289810) to evaluate the superiority of tiragolumab plus atezolizumab over atezolizumab in treatment-naïve patients with advanced NSCLC expressing PD-L1; however, the recently released results of the interim analysis showed that the trial did not meet the PFS; hence, the study will continue investigating the OS until the next planned analysis (99). Currently, other phase III studies of anti-TIGIT antibodies are underway, and the competition to develop anti-TIGIT antibodies is starting to intensify. Phase III studies on domvanalimab combined with zimberelimab, an anti-PD-1 antibody (ARC-10 study, NCT04736173), and durvalumab as maintenance therapy after chemoradiotherapy (PACIFIC-8, NCT05211895) are ongoing, while the efficacy of adding vibostolimab to pembrolizumab or pembrolizumab plus platinum combination chemotherapy as first-line treatment is being examined in phase III trials (NCT04738487, NCT05226598) based on the promising results of a phase I study (100).

2.2.4 Angiogenesis inhibitors

Following the success of the IMpower150 trial on bevacizumab in addition to platinum-based chemotherapy and atezolizumab as first-line treatments for patients with advanced NSCLC, a number of clinical trials have been conducted combining angiogenesis inhibitors and ICIs (101). A placebo-controlled phase III trial (ONO-4538-52/TASUKI-52) was conducted to evaluate the superiority of nivolumab plus carboplatin plus paclitaxel plus bevacizumab over carboplatin plus paclitaxel plus bevacizumab in patients with advanced non-squamous NSCLC and PD-L1 expression of ≥1%; the median PFS times were 12.1 months (96.37% CI, 9.8–14.0) and 8.1 months (96.37% CI, 7.0–8.5), respectively, with an HR of 0.56 (96.4% CI, 0.43–0.71; p < 0.0001), with significantly better outcomes in the nivolumab arm regardless of tumor PD-L1 expression status (102). The median OS was similar, but the HR showed a favorable trend, indicating that this combination strategy could be a potential first-line treatment for non-squamous NSCLC. A multicenter, open-label, single-arm phase II study (@Be trial) that evaluated the efficacy and safety of atezolizumab plus bevacizumab as first-line therapy in 39 non-squamous NSCLC patients with a PD-L1 tumor proportion score of ≥50% was conducted by the West Japan Oncology Group, and the results were reported at the 2020 European Society for Medical Oncology conference (103). The ORR was 64.1%, with tumor shrinkage observed in most patients, and the safety was comparable to the previously reported data. The West Japan Oncology Group is planning to conduct a @Be-F1rst
ICI resistance is being considered.

possible use of this regimen in the front-line setting to overcome therapy, showing an ORR of 10% with acceptable safety. The after treatment with ICIs, were treated with this combination safety of combining BI 836880, a bispecific

adding bevacizumab. Another phase I trial investigating the response, suggesting that ICI resistance was overcome by angiopoietin-2 in addition to vascular endothelial growth factor, and ezabenlimab, an anti-PD-1 antibody, has been reported (105). Forty patients with NSCLC, whose disease had progressed and received atezolizumab until the detection of disease progression following atezolizumab monotherapy (104). This trial enrolled ICI-naive pretreated NSCLC patients whose disease progressed after at least one line of platinum-based chemotherapy. Patients received atezolizumab until the detection of disease progression on radiographic evaluation (stage I, n = 42). Bevacizumab was combined with atezolizumab (stage II, n = 24). The disease control rate in patients with stage II disease was 87.5% (95% CI, 67.6–97.3) including 12.5% of those who achieved partial response, suggesting that ICI resistance was overcome by adding bevacizumab. Another phase I trial investigating the safety of combining BI 836880, a bispecific nanobody targeting angiopoietin-2 in addition to vascular endothelial growth factor, and ezabenlimab, an anti-PD-1 antibody, has been reported (105). Forty patients with NSCLC, whose disease had progressed after treatment with ICIs, were treated with this combination therapy, showing an ORR of 10% with acceptable safety. The possible use of this regimen in the front-line setting to overcome ICI resistance is being considered.

2.2.5 TIM-3 inhibitors

Although it lags behind LAG-3 and TIGIT, anti-T-cell immunoglobulin and mucin domain 3 (TIM-3) therapy is expected to be developed because the co-expression of TIM-3 and PD-L1 adversely affects the immune system and is effective as a combination therapy (106). In 2018, the results of a phase I trial of a combination of cobolimab, an anti-TIM-3 antibody, and dostarlimab, an anti-PD-1 antibody, showed that in 25 NSCLC patients who developed resistance to anti-PD-1 antibodies, cobolimab combined with dostarlimab produced a response in 3 of 20 evaluable patients; hence, further development of this combination therapy is expected to overcome ICI resistance (107). A phase II/III trial (COSTAR Lung Study, NCT04655976) is currently performed to evaluate the superiority of cobolimab plus dostarlimab plus docetaxel and dostarlimab plus docetaxel over docetaxel alone in NSCLC, with an expected completion date of 2024. Meanwhile, sabatolimab, an anti-TIM-3 antibody, was granted a fast-track designation by the FDA and Orphan Medical Product designation by the European Medicines Agency for the treatment of myelodysplastic syndrome based on the results of a phase I trial on myelodysplastic syndrome and acute myeloid leukemia presented at the 2019 American Society of Hematology conference (108). However, for solid tumors, the combination of cobolimab and anti-PD-1 antibody spartalizumab yielded a response rate of only 6% as reported in a phase I trial; another phase I trial showed a modest response rate of 4% for LY3321367 (an anti-TIM-3 antibody) combined with LY300054 (an anti-PD-L1 antibody), making it difficult to decide the necessity of developing anti-TIM-3 therapies in the future (109, 110).

2.3 Other anti-PD-1/PD-L1 antibody combination therapies

2.3.1 Lenvatinib

Combination therapy with the multi-kinase inhibitors lenvatinib and pembrolizumab has been approved by the FDA, European Medicines Agency (EMA), and Pharmaceuticals and Medical Devices Agency (PMDA) for the treatment of uterine cancer and/or renal cell carcinoma. Basic studies have shown that the combination of lenvatinib and pembrolizumab improves the immune microenvironment in hepatocellular carcinoma, and several clinical trials are underway to expand its indication to other types of cancer (111, 112). Several phase III trials using lenvatinib in NSCLC are being examined the efficacy of pembrolizumab in combination with pemetrexed plus platinum in the first-line treatment of non-squamous NSCLC patients (LEAP-006, NCT03829319), in combination with pembrolizumab in the first-line treatment of PD-L1-positive patients (LEAP-007, NCT03829332), in combination with docetaxel in the second-line setting (LEAP-008, NCT03976375), and in combination with KN046, a recombinant humanized PD-L1/CTLA-4 bispecific fusion protein, after ICI resistance (NCT05001724). Ongoing phase I and II trials are using lenvatinib combination therapy with pembrolizumab in the perioperative treatment (NCT04875585), pembrolizumab plus pemetrexed plus carboplatin following treatment with epidermal growth factor receptor-tyrosine kinase inhibitors in epidermal growth factor receptor mutation-positive patients with NSCLC (NCT05258279), GI-101, a bispecific fusion protein of CD80 and interleukin (IL)-2 mutants (NCT04977453), IRB318, a bispecific antibody of PD-1 and PD-L1 (NCT04777084), and envalolimab, a subcutaneous anti-PD-1 antibody formulation (NCT05024214). It might not be long before lenvatinib becomes available in the clinical setting for patients with thoracic malignancies following the approval for thymic cancer in Japan (113).

2.3.2 Canakinumab

The inflammatory cytokine IL-1β is believed to be involved in cancer invasion, progression, and metastasis. A subgroup analysis of the CANTOS trial showed that canakinumab, an IL-1β inhibitor with an anti-inflammatory effect on atherosclerosis that inhibits the recurrence of myocardial infarction, also reduces the incidence of lung cancer and death (114). Based
on these findings, the efficacy of canakinumab in NSCLC was validated by multiple phase III trials; however, the addition of canakinumab to standard therapy as first-line (CANOPY-I) and second-line or later-line (CANOPY-II) treatments failed to meet the primary endpoint, as reported in 2021 (115, 116). Canakinumab is currently under a phase II trial as a preoperative treatment (CANOPY-N, NCT03968419) and a phase III trial as a postoperative treatment (CANOPY-A, NCT03447769), being expected to improve the immune microenvironment through IL-1β inhibition in early stage NSCLC.

2.3.3 Other novel immunotherapy agents

Other new agents in the early stages of development for combination therapy with ICIs include MK-4830, an IgG4 monoclonal antibody that targets the immunoglobulin-like transcript 4 receptor. The preliminary results of a phase I trial, which was first presented at the 2020 European Society for Medical Oncology conference and published in 2021, showed a promising ORR of 24% in a dose-escalation cohort treated with pembrolizumab; notably, five of the eleven patients who were resistant to anti-PD-1/PD-L1 antibody therapy had an objective response (117). The efficacy and safety of MK-4830 are verified further in a substudy of the phase II KEYMARKER trial (NCT04165083 and NCT04165096). MK-5890, an anti-CD27 agonist, has also been evaluated in the KEYMARKER trial; according to the phase I results presented at the 2019 Society for Immunotherapy of Cancer conference, combination therapy with pembrolizumab showed an ORR of 10.5%. Of the 14 patients who switched to combination therapy with pembrolizumab after experiencing disease progression following MK-5890 monotherapy, 5 patients, including 2 who achieved complete response, showed a favorable response (ORR, 35.7%), which was a very promising result (118).

3 Future perspective

PD-1/PD-L1 antibodies are generally approved for NSCLC and can be used in clinical practice in Japan as well as in the US and EU, except for cemiplimab. However, in solid tumors other than NSCLC, pembrolizumab for cervical cancer was approved by the FDA in 2018, cemiplimab and pembrolizumab for cSCC in 2018 and 2020, respectively, and cemiplimab for cBCC in 2021; however, none of these agents were approved for in Japan because Japanese patients were not enrolled in pivotal trials (119–121). In addition, a number of PD-1/PD-L1 antibodies developed in China have not been approved by the PMDA, FDA, or EMA, which is thought to be due in part to the rise of emerging biopharma companies (EBPs)—those with an estimated expenditure on research and development (R&D) of less than $200 million and less than $500 million in revenue—which contributed more than 70% of the FDA regulatory submission for approval (122, 123). Most EBPs are located in the US, EU, or China and do not own any Japanese corporations or domestic administrators, which has led to an increasing number of cases that have not been conducted in Japan. This “decentralized drug development” in oncology field is expected to expand and accelerate in the near future, and certain drugs that can be used in other countries might not be available in our own countries. These problems are so complex that they are difficult to solve without establishing relevant policies (124).

In the past two decades, “drug lag” (i.e., the delay in time required for the approval of oncology drugs) was an issue in Japan compared with that in the US or EU. Efforts have been made to eliminate the “drug lag” with other countries using several approaches such as expediting the regulatory approval, establishment of the Strategy of SAKIGAKE, which allows the accelerated approval of drugs as breakthrough therapies and addressing unmet medical needs in Japan attracting foreign drug trials including orphan drugs, and launching of the Advanced Medical Care Program to enable patients to gain access to promising unapproved drugs or medical devices through the National Health Insurance (NHI) coverage (125–129). In fact, review periods for new drugs have been shortened, with the median periods in 2018 being 10.0 months for the PMDA, 10.4 months for the FDA, and 13.7 months for the EMA (130).

Although the review period gap with overseas countries is beginning to shorten, the number of unapproved drugs in Japan has reached 70% due to the rise of EBPs and the negative impact of the expected shrinking of the Japanese pharmaceutical market and the biennial revision of Japanese drug prices (122, 129–132). Currently, although no specific national policies or measures that can attract foreign EBPs to Japan are prominently practiced, our institution has opened its doors to domestic and foreign entrepreneurial ventures, providing online consultation by experts with experience in regulatory review and clinical trial initiatives (133). In addition to promoting the understanding of Japan’s strengths, such as NHI coverage, which covers the medical expenses for treatment and medical testing, and Japan’s market value (once approved, the drug always carries a drug price and is likely to be delivered to patients), further political support is needed to promote drug development in Japan, such as encouraging more Japanese participation in international joint clinical trials (134–136). To maximize the benefits of “decentralized drug development” for Japanese patients, the Japanese industry, government, and academia should cooperate and catch up with the global market.

4 Discussion

The ICI therapy has drastically rewritten the history of cancer treatment. The development of novel agents, and use of combination therapies will continue to accelerate cancer
treatment. In addition to the clinical trials reported to date, numerous drug combinations, such as antibody-drug conjugates, bispecific antibodies targeting multiple immune-checkpoint molecules, and other immune-related molecules in combination with ICIs, have been used. Although several compounds for immunotherapy are being developed, the effectiveness of immunotherapy often depends on the immune and systemic status of the host, and results from early phase trials are not often reflected in later phase trials. Although some treatments are effective in certain patients, there is also a need for measures to address the disadvantages associated with such treatments, such as the increasing medical costs, the need to implement scientific approaches to narrow down the population that will benefit from the treatment (or identifying those who will not benefit), and the duration of treatment for patients who do benefit. In a randomized post-marketing trial of NSCLC patients (CheckMate-153 study), the PFS and OS were significantly shorter after nivolumab discontinuation (137). However, the impact of discontinuation cannot be concluded in this trial alone as it was not planned to address a specific statistical hypothesis: patients who showed exacerbation were included, the number of patients who responded to nivolumab was much larger in the continuation arm, and the important patient characteristics were imbalanced between the treatment arms. The Japan Clinical Oncology Group is currently conducting the JCOG1701 trial (SAVE study), a randomized controlled phase III trial that aimed to evaluate the non-inferiority of treatment suspension to continued treatment in NSCLC patients who have benefited from anti-PD-1/PD-L1 antibody treatment for at least 12 months (138). The prognostic and predictive roles of circulating tumor DNA will be explored in this study. Another randomized phase II–III trial for NSCLC compared the outcomes of discontinuation and continuation of pembrolizumab after six months of treatment with chemotherapy plus pembrolizumab combination therapy (139). These trials will elucidate the optimal management of NSCLC patients treated with ICIs from the perspective of not only safety and better biomarkers, but also cost-effectiveness.

As it has become common for multiple departments to collaborate in the treatment of immune-related adverse events, further development of strategies is warranted to provide optimal treatment to cancer patients through a global collaboration between industry, the government, and academia worldwide.

Author contributions

This review was drafted by TM and NY, and critically revised by YK, JS, TK, and TS. All authors contributed to the article and approved the submitted version.

Conflict of interest

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The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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