Immune checkpoint inhibitors for small cell lung cancer: opportunities and challenges

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Abstract: Lung cancer is the most common cancer and the leading cause of cancer death worldwide, with an estimated 2.1 million new cases and 1.8 million deaths in 2018. Although small cell lung cancer (SCLC) is the most aggressive type of lung cancer, it shows high response rates to chemotherapy in early lines of therapy. Unfortunately, it is associated with rapid recurrence and relatively poor prognosis. Over the last few years, considerable progress has been made in cancer immunotherapy. One of the most promising ways to activate therapeutic antitumor immunity is via blockade of immune checkpoints, such as cytotoxic T lymphocyte-associated protein-4 (CTLA-4) and programmed cell death protein-1/programmed cell death ligand-1 (PD-1/PD-L1). Immune checkpoint inhibitors show promise as SCLC therapeutics. The overall expectation for immuno-oncology is high, and the outcomes of trials will hopefully reveal a variety of treatment options for SCLC patients. In this review, we discuss the discovery of new immune inhibitory and stimulatory pathways and rational combination strategies to explain the role of immunotherapy in SCLC and its future opportunities and challenges.

Keywords: small cell lung cancer, immune checkpoint inhibitors, CTLA4, cytotoxic T lymphocyte-associated protein4, PD1, programmed cell death protein1, PDL1, programmed cell death ligand 1, next generation of immune checkpoints

Introduction

Lung cancer is the most common cancer and the leading cause of cancer death worldwide, with an estimated 2.1 million new cases and 1.8 million deaths in 2018.1 Approximately 13% of lung cancer patients have small cell lung cancer (SCLC).2 Although this aggressive tumor shows high response rates to chemotherapy in early lines of therapy, it is associated with rapid recurrence and relatively poor prognosis and is often diagnosed at late stages with systemic metastasis.3,4 Thus, the 5-year survival of SCLC patients is very low and varies according to stage, with 5-year relative survival rates of 31% for limited-stage SCLC (LS-SCLC) but just 2% for extensive-stage SCLC (ES-SCLC).5

Because SCLC is generally highly sensitive to chemotherapy and radiation therapy, combined chemotherapy and radiotherapy has become the accepted standard treatment for all stages of SCLC. The standard chemotherapy regimen of cisplatin or carboplatin plus etoposide used as the first-line treatment of LS-SCLC and ES-SCLC diseases has not changed over the past four decades. Radiotherapy is administered to those patients with LS-SCLC whose cancer is confined to the chest in a single tolerable radiation field.6-8 Recently, the randomized phase III CONVERT trial9 was designed to show the superiority of once-daily concurrent...
Recently, the results of the IMpower study showed that the combination of atezolizumab (TECENTRIQ®) with carboplatin and etoposide chemotherapy for the initial treatment of ES-SCLC (median OS [mOS] of 12.3 months in the atezolizumab group and 10.3 months in the placebo group; hazard ratio [HR] 0.70, 95% confidence interval [CI] 0.54–0.91, p=0.007). In Figure 1, we summarize the standard of care therapies and new therapies that have been approved by the FDA.

Additionally, the FDA has granted a priority review designation to a supplemental biologics license application (sBLA) for pembrolizumab (KEYTRUDA®) as a treatment for patients with advanced SCLC whose disease has progressed after two or more prior lines of therapy. The application is based on findings from cohorts of the phase II KEYNOTE-158 and phase Ib KEYNOTE-028 studies, in which pembrolizumab elicited 19% and 33% overall response rates (ORRs) in patients with advanced and ES-SCLC, respectively. The immune system can recognize physiological and pathological changes at the cellular level. After tumorogenesis, tumor-associated antigens can be presented by antigen-presenting cells (APCs) to major histocompatibility complex (MHC) I and then recognized by T cell receptor on CD8 cytotoxic T cells. T cell activation is induced by a secondary co-stimulatory signal, namely, the binding of B7 protein on APCs to CD28 on cytotoxic T cells. Afterward, the activated T cell induces the death of the physiologically and pathologically altered cell. Conversely, the activated T cells are negatively regulated by increased expressed of cytotoxic T lymphocyte-associated protein 4 (CTLA-4), which has a higher affinity for B7 protein of APCs than the CD28 molecule of cytotoxic T cells. CTLA-4 is normally expressed in regulatory T cells (Tregs) and acts as a restraint molecule of the activated immune system, suppressing auto-reactivity. Similar to the CTLA-4 molecule, PD-1 is an immune checkpoint inhibitor in activated T cells, peripheral B cells, and myeloid cells. In contrast, the binding of PD-1 to PD-L1 on tissues or hematopoietic cells occurs in the late phase and thereby mainly suppresses activated T cells in peripheral immune effector sites rather than in central lymph systems. Overexpression of tumor-specific antigens on tumor cells can facilitate their recognition by the host immune system, especially by T cells. CTLA-4 and PD-1 are recognized as inhibitors of immune effector cells and they therefore contribute to the rapid development of cancer cells due to inactivation of the immune system during tumorigenesis.

Figure 1 Timeline of treatment for SCLC. This timeline illustrates the standard of care therapies and new therapies that have been approved by the FDA.

Radiotherapy
Chemotherapy
Immunotherapy

2019
2018
2007
1999
1996
1992
1985

Atezolizumab
Nivolumab (metastatic SCLC)
Topotecan (relapsed SCLC), (oral) 2007
PCI (LD-SCLC) 1999
45Gy b.i.d. (LS-SCLC) 1999
Carboplatin+etoposide 1999
Topotecan (relapsed SCLC), (injection) 1996
Thoracic radiotherapy (LS-SCLC) 1992
Cisplatin+etoposide 1985

Timeline of treatment for SCLC. This timeline illustrates the standard of care therapies and new therapies that have been approved by the FDA.
Because SCLC has the highest rate of tumor mutational burden (TMB), the tumor should be sensitive to immune-based therapeutic approaches. TMB is used as a predictor of immune checkpoint inhibitor efficacy for many cancer types. In Helmann et al, the objective response rate was 21.3% in the high TMB group (248 mutations on whole-exome sequencing) but just 4.8% in the lower TMB group (0 to 142 mutations). PD-L1 expression has been applied as a biomarker to predict the efficacy of PD-1/PD-L1 inhibitors in relation to the dynamic and heterogeneous expression of PD-L1 in the tumor microenvironment. Immune checkpoint molecules are frequently overexpressed in SCLC-associated tumor-infiltrating lymphocytes. Figure 2 summarizes the inhibitory and stimulatory molecules and their targets. Cancer immunotherapy research achievements can represent a new hope for SCLC patients and confer them with improved and sustainable survival opportunities. Thus, in this review, we aim to provide an up-to-date overview of recent developments in immune checkpoint therapy and novel immune checkpoint agents and combination therapies in SCLC, focusing on clinical trials.

Figure 2. Immunosuppressive and immunostimulatory checkpoints and their targets. T cells recognize antigens presented by MHC complex on APCs or tumor cells through T cell receptor (TCR). Many of the ligands bind to multiple receptors; some deliver co-stimulatory signals, whereas others deliver inhibitory signals. Full activation of T cells is mediated by binding of CD28 to B7. CTLA-4 competes with CD28 for binding to B7 and transmits an inhibitory signal suppressing T cell activation. PD-1 is expressed on T and NK cells after antigen exposure. Following binding of PD-1 to its ligands, such as PD-L1/PD/L2 (expressed on tumor cells or APCs), an inhibitory signal turns off the immune response mediated by activated T and NK cells. A2aR is expressed on T and NK cells, similar to PD-1. As in CTLA-4 and PD-1 pathways, a significant influence of A2aR signaling on Tregs and effector T cells is likely to be the fundamental driving force behind its immunosuppressive effect. TIM3 is an inhibitory receptor expressed by T and NK cells that interacts with soluble or cell surface galectin 9 to negatively regulate T and NK cell function. BTLA is an inhibitory receptor that is structurally and functionally related to CTLA-4 and PD-1 and expressed by T cells. LAG3 is a receptor that forwards inhibitory signals to activated T and NK cells through LAG3/TCR-MHC interaction, leading to downregulation of immune responses. KIR receptors are expressed on NK cells and to a lesser extent on CD8+ T cells. The binding of inhibitory KIRs (inhKIRs) to their ligands (HLA class I molecules) protects normal cells against the NK cell attack. In contrast, in cancer, higher expression of inhKIRs on the surface of NK and T cells promotes a downregulation of their function. OX40 is a stimulatory receptor, highly expressed by activated T cells and to a lesser degree by NK cells. OX40 and its ligand OX40L play a pivotal role in the activation, potentiation, proliferation, and survival of T cells and in modulating NK cell function. CD40 is a stimulatory receptor expressed on T cells. The interaction between CD40 and CD154 stimulates cytokine secretion by B cells with subsequent T cell activation and tumor cell death. The levels of ICOS, a specific T cell co-stimulatory receptor expressed on T cells, are upregulated in activated T cells, particularly after the use of anti-CTLA-4 therapies. CD27 is a stimulatory receptor expressed on T cells. CD27–CD70 binding results in a potent signal that activates and differentiates T cells into effector and memory cells and boosts the level of B cells. GITR is a co-stimulatory receptor expressed on T and NK cells. 4-1BB is an inducible co-stimulatory receptor expressed on T and NK cells. The interaction between 4-1BB and 4-1BBL triggers subsequent immune cell proliferation and activation, particularly of T and NK cells. TIGIT is a member of CD28 family-like receptors expressed on T and NK cells. It exerts direct immunosuppressive effects on these cells and indirectly increases the release of immunoregulatory cytokines, decreases the production of interferon-γ and IL-7, and prevents maturation of dendritic cells.
The CTLA-4 pathway and CTLA-4 checkpoint inhibitors

CTLA-4 was the first immune checkpoint receptor to be targeted by a therapeutic agent. Monoclonal antibodies against CTLA-4 (ipilimumab and tremelimumab) are designed to prevent the interaction between CTLA-4 and its ligands (CD80/CD86). In this manner, they blockade the inhibitory signal provided by CTLA-4 and enhance the activation and proliferation of tumor-specific T cells, facilitating an effective immune response against tumor cells. In 2011, ipilimumab became the first immune checkpoint inhibitor to be approved by the FDA.

Ipilimumab has achieved promising results in SCLC treatment. In a three-arm phase II trial (NCT00527735), patients received either six cycles of carboplatin and paclitaxel with or without ipilimumab or with ipilimumab in a phased schedule. The immune-related progression-free survival (irPFS) was significantly prolonged for the phased ipilimumab arm (6.4 vs. 5.3 months, HR 0.64, P=0.03) but was not prolonged for the concurrent arm compared with the control (5.7 vs. 5.3 months, HR 0.75, P=0.11). There was also a nonsignificant trend for improved OS in the phased ipilimumab arm (12.5 months, 95% CI 7.9–14.9), compared with mOSs of 9.1 months (95% CI 6.7–13.0) for the concurrent arm and 10.5 months (95% CI 8.6–11.7) for the control arm. The HR values relative to control were 0.76 (95% CI 0.48–1.19, P=0.13) for phased ipilimumab and 0.89 (95% CI 0.57–1.39, P=0.41) for concurrent ipilimumab. There was also increased toxicity, with grade III and IV toxicities more common in the ipilimumab arms (30% concurrent vs. 43% concurrent vs. 50% phased). The study did not identify any potential biomarkers that could predict the response to ipilimumab treatment in patients with SCLC.

In an open-label phase II study of ipilimumab with carboplatin/etoposide for use as first-line treatment in previously untreated patients with ES-SCLC, patients received carboplatin and etoposide up to six cycles (3–6) with ipilimumab. The presence of autoantibodies (anti-SOX2, anti-HU, anti-Yo, anti-VGCCA, anti-VGPCA, thyroid peroxidase, rheumatoid factors, anti-muscle antibodies, ANA, and ANCA) was investigated at baseline in 38 patients. Ipilimumab in combination with carboplatin and etoposide might benefit a subgroup of patients with advanced SCLC. A positive autoimmune profile at baseline was associated with improved outcomes, with patients with any positive autoantibodies at baseline having a significantly longer median irPFS (8.8 months [95% CI 5.1–10.7] vs. 7.3 months [95% CI 2.9–7.9], P=0.036). Antinuclear antibody positivity predicted a significantly prolonged irPFS (10.2 months vs. 6.9 months, P=0.032). The mPFS (mPFS) was 6.9 months (95% CI 5.5–7.9), the median irPFS was 7.3 months (95% CI 5.5–8.8), and the mOS was 17 months (95% CI 7.9–24.3). Increased toxicity was detected, with grade III or higher toxicity developing in 90%; 69% of these toxicities were thought to be related to ipilimumab.

Moreover, in a double-blind phase III study of ipilimumab plus etoposide and a platinum agent, OS was not increased compared with chemotherapy alone (mOS of 11.0 vs. 10.9 months, HR 0.94, 95% CI 0.81–1.09, P=0.3775). There was also no significant improvement in the mPFS (4.6 vs.4.4 months; HR 0.85, 95% CI 0.75–0.97). Rates of treatment-related adverse events were 48% for grade III to IV events in the ipilimumab arm and 44% in the placebo arm (Table 1).

PD-1 and PD-L1 pathways and PD-1 and PD-L1 checkpoint inhibitors

The PD-1 receptor is another important checkpoint receptor being studied in oncology. In contrast to CTLA-4, PD-1 is expressed on a broader range of immune cells besides T lymphocytes (e.g., B lymphocytes and natural killer [NK] cells). Antibodies designed to inhibit the interaction between PD-1 and its ligands (PD-L1 and PD-L2) have shown promising results in clinical trials of different tumor types, including SCLC. A monoclonal anti-PD-1 antibody (nivolumab) was designated a breakthrough therapy for the treatment of patients with metastatic SCLC on August 17th, 2018. The approval was based on the durability of responses in the phase I/II CheckMate-032 trial, in which the objective response rate was 12% (95% CI 6.5–19.5) for nivolumab after platinum-based chemotherapy and one other prior line of therapy in 109 patients, according to the new label. This response rate comprised 11% partial responses and 0.9% complete responses. The median duration of response was 17.9 months (95% CI 7.9–42.1), with 62% of patients continuing to respond at 12 months and 39% still responding at 18 months.

In the CheckMate-032 study, the mOS was 4.4 months (95% CI 3.0–9.3) in the nivolumab monotherapy arm but 6.0 months (95% CI 3.6–11.0) in the nivolumab 3 mg/kg plus ipilimumab 1 mg/kg (N3/11) arm and
<table>
<thead>
<tr>
<th>Agent</th>
<th>Trials identifier</th>
<th>Phase</th>
<th>Study design</th>
<th>Patients</th>
<th>Primary endpoint</th>
<th>Outcome</th>
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<td>130</td>
<td>irPFS</td>
<td>irPFS 6.4, 5.7, 5.3 mPFS 5.2, 3.9, 5.2 mOS 12.5, 9.1, 10.5 (phased, concurrent and control respectively)</td>
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<td></td>
<td>NCT01331525</td>
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<td>Open label single group assignment: Ipilimumab+Carboplatin+Etoposide</td>
<td>42 (38)</td>
<td>PFS</td>
<td>mPFS 6.9 (95% CI, 5.5–7.9); median irPFS 7.3 (95% CI, 5.5–8.8); mOS 17 (95% CI, 7.9–24.3)</td>
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<td>NCT01450761</td>
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<td>Randomized, Multicenter, Double-Blind: Arm A: Ipilimumab+ET+Cisplatin +Carboplatin; Arm B: Placebo + ET+Cisplatin +Carboplatin</td>
<td>1132</td>
<td>OS</td>
<td>OS 11.0 vs 10.9 months, HR=0.95 (95%CI 0.8–1.09 p=0.3775), PFS 4.6 vs 4.4 months HR=0.85(95%CI 0.75–0.97); (treatment vs placebo)</td>
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<td>RR</td>
<td>mPFS 1.4 (95%CI, 1.3–2.8) mOS 9.6 (95%CI, 7.0–12)</td>
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<td>CheckMate032</td>
<td>Randomized, open label, parallel assignment: Arm A: Nivolumab monotherapy Arm B: Nivolumab+Ipilimumab</td>
<td>1150</td>
<td>ORR</td>
<td>ORR 11.9% mPFS 1.4 months (95% CI, 1.3–1.6) mOS 5.6 months (95% CI, 3.1–6.8) 1-year OS 28.3% (95% CI, 20.0–37.2).</td>
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</table>

Abbreviations: SCLC, small cell lung cancer; ET, etoposide; irPFS, immune-related progression-free survival; PFS, progression-free survival mOS, median overall survival; OS, overall survival; RR, risk ratio; ORR, objective response rate; HR, hazard ratio; CI, confidence interval.
7.7 months (95% CI 3.6–18.0) in the nivolumab 1 mg/kg plus ipilimumab 3 mg/kg (N1/I3) arm. The mPFS was 1.4 months in the nivolumab monotherapy arm (95% CI 1.4–1.9) and 1.4 months (95% CI 1.3–2.2) and 2.6 months (95% CI 1.4–4.1) in the N3/I1 and N1/I3 arms, respectively. The 1-year PFS was 11% (95% CI 5.0–19.0) in the nivolumab monotherapy arm. An updated result of the CheckMate-032 study was recently published.\(^{39}\) Nivolumab monotherapy against SCLC had a strong therapeutic effect and was well tolerated as a third- or later-line treatment for recurrent SCLC. The objective response rate was determined to be 11.9% in the patients with recurrent LS- or ES-SCLC, the mPFS was 1.4 months (95% CI 1.3–1.6), and the mOS was 5.6 months (95% CI 3.1–6.8). Moreover, the 1-year OS was 28.3% (95% CI 20.0–37.2). However, the open-label Checkmate-331 study failed to meet the primary endpoint of OS compared with standard of care.\(^{10}\)

A randomized, open-label, phase I/II study comparing nivolumab with 177Lu-DOTA0-Tyr3-Octreate radiation therapy, sequentially, in patients with ED-SCLC is currently active and not recruiting patients (NCT03325816).\(^{44}\) The primary endpoints are PFS and the maximum tolerated dose of 177Lu-DOTA0-Tyr3-Octreate. Nivolumab 240 mg is being administered every 2 weeks until disease progression, with 177Lu-DOTA0-Tyr3-Octreate administered 2 weeks after the first administration of nivolumab. Each dose is infused over 30 minutes. A randomized, open-label, phase II study comparing nivolumab treatment with no treatment after chemotherapy in ED-SCLC is also currently active with the primary endpoint of PFS (NCT03382561).\(^{44}\) In addition, nivolumab clinical trials are ongoing and recruiting patients, including an open-label, phase II study of patients with advanced SCLC treated with gemcitabine (NCT03662074) (Table 2).\(^{44}\)

### Pembrolizumab

Pembrolizumab (KEYTRUDA\(^{®}\)), a fully humanized IgG4 isotype antibody against PD-1, is recognized as an effective immunotherapy for several solid tumors. The FDA initially approved it to treat metastatic melanoma and further approved it in 2017 for any unresectable or metastatic solid tumor with certain genomic anomalies.\(^{40}\) A multicenter phase II trial\(^{41}\) of maintenance pembrolizumab in 45 patients with SCLC showed no improvement in the mPFS (1.4 months, 95% CI 1.3–2.8) when patients received pembrolizumab after completion of 4 to 6 cycles of platinum/etoposide, with a 1-year PFS of 13%. The mOS was 9.6 months (95% CI 7.0–12.0) and the 1-year survival was 37%. Further analysis indicated that the mPFS was 6.5 months (95% CI 1.1–12.8) in the PD-L1-positive group and 1.3 months (95% CI 0.6–2.5) in the PD-L1-negative group, suggesting that this agent only benefits a small subset of PD-L1-expressing patients. The phase 1b basket study KEYNOTE-028 and the phase II basket study KEYNOTE-158 evaluated the antitumor activity of pembrolizumab in patients with advanced SCLC who had received at least two lines of prior therapy,\(^{42}\) with pembrolizumab demonstrating promising antitumor activity. In this exploratory pooled analysis, the mOS was 7.7 months and the 1- and 2-year survival rates were 34.3% and 20.7%, respectively. The mPFS was 2 months, and the 1- and 2-year survival rates were 16.9% and 13.1%, respectively. Moreover, several ongoing clinical trials are evaluating the efficacy and safety of pembrolizumab in SCLC, including a phase III study of patients with ED-SCLC treated with etoposide/platinum (cisplatin or carboplatin) in combination with pembrolizumab (Keynote-604: NCT03066778), as well as in combination with chemotherapy or chemoradiotherapy (Table 2). In addition, the REACTION trial,\(^{44}\) with a primary endpoint of PFS, is evaluating platinum/etoposide in treatment naïve patients with or without pembrolizumab.

### Atezolizumab

Atezolizumab (MPDL3280A), a fully humanized engineered IgG1 monoclonal antibody against PD-L1, is an effective immunotherapy for several solid cancers. Recently, the FDA granted approval to the combination of atezolizumab (TECENTRIQ\(^{®}\)) with carboplatin and etoposide for the frontline treatment of patients with ES-SCLC. The approval is based on findings from the phase III IMpower133 study,\(^{11}\) which demonstrated that the addition of atezolizumab to carboplatin and etoposide significantly improved OS in patients with ES-SCLC. The mOS was 12.3 months in the atezolizumab group and 10.3 months in the placebo group (HR 0.70, 95% CI 0.54–0.91, P=0.007. The mPFS was 5.2 months and 4.3 months, respectively (HR 0.77, 95%CI 0.62–0.96, p=0.02).\(^{12}\) However, a randomized phase II trial (IFCT-1603)\(^{43}\) investigating the therapeutic efficacy of single atezolizumab for relapsed SCLC revealed that atezolizumab did not have a significantly better therapeutic efficacy and safety profile. PFS was determined to be 1.4 months in
**Table 2** Ongoing trials of immune checkpoint agents in SCLC

<table>
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<tr>
<th>Agent</th>
<th>Trials identifier</th>
<th>Phase</th>
<th>Study design</th>
<th>Patient, estimated</th>
<th>Estimated Study completion Date</th>
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<td>Phase I: Confirmation of Recommended Phase II Dose; Phase II: PFS</td>
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<td>60</td>
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<td>II</td>
<td>NCT03066778</td>
<td>Registered, multicenter, double-blind, parallel assignment: Arm A. Pembrolizumab+EP Arm B. Placebo+EP</td>
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<td>Laboratory findings, AEs</td>
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<td>Non-randomized, open label, parallel assignment: for SCLC: Arm A: Durvalumab+CT+SRT Arm B: Durvalumab+CT+HRT hyperfractionated Arm C: Durvalumab+Tremelimumab+CT +SRT Arm D: Durvalumab+Tremelimumab+CT +HRT</td>
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<td>ORR</td>
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</table>
the atezolizumab group (95% CI 1.2–1.5) but 4.2 months in the chemotherapy group (95% CI 1.5–5.9). In this field of study, several clinical trials are still being performed to evaluate the efficacy and safety of atezolizumab for SCLC. In addition, other PD-L1 immune checkpoint inhibitors—durvalumab and avelumab—are also being investigated in clinical trials.44

**Combination therapy**

A number of ongoing trials are focused on combination therapies, such as a phase I/II trial investigating ROV A-T (an anti-DLL3 antibody drug conjugate) with nivolumab and ipilimumab (NCT03026166) and a phase I/II trial investigating olaparib (PARP inhibitor) with durvalumab (anti-PD-L1 antibody) and the anti-VEGF antibody bevacizumab (NCT02734004). A randomized, open-label, phase II study (STIMULI) comparing ipilimumab following chemoradiotherapy alone in LS-SCLC is currently recruiting patients (NCT02046733) with primary endpoints of OS and PFS. In the induction phase, nivolumab is to be administered at a dose of 1 mg/kg i.v., followed by ipilimumab 3 mg/kg i.v. once every 3 weeks for four cycles. Nivolumab will be continued as maintenance therapy at a dose of 240 mg i.v. once every 2 weeks for a maximum of 12 months; the results are expected in 2022. Phase III data from the CheckMate-451 trial showed that OPDIVO® (nivolumab) combined with YERVOY® (ipilimumab) missed the primary endpoint of OS.45 Preliminary results from the BALTIC study,46 which evaluated the combination of durvalumab and tremelimumab in 25 patients with platinum-refractory/platinum-resistant ES-SCLC, showed an objective response rate of 9.5%, mPFS of 1.9 months, and mOS of 6.0 months, suggesting improved activity over single-agent anti-PD-L1 therapy (Table 2).

A phase I/II study44 comparing nivolumab+ipilimumab ±plinabulin in recurrent SCLC is currently recruiting patients (NCT03575793). Primary endpoints include the maximum tolerated dose and PFS. In the phase I trial, nivolumab is to be intravenously administered at a dose of 1 mg/kg i.v., followed by ipilimumab 3 mg/kg i.v. and plinabulin in dose-escalating cohorts (level 1, 13.5 mg/m²; level 1 (start), 20 mg/m²; level 2, 30 mg/m²). After four treatment cycles, ipilimumab will be stopped and patients will be administered nivolumab 240 mg and plinabulin every 2 weeks; the results are due in 2022. A number of ongoing trials are focusing on durvalumab and tremelimumab combination therapies, such as the CASPIAN and CLOVER trials. A phase III study (CASPIAN) comparing

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</table>
| Table 2 (Continued).

**Abbreviations:** SCLC, small cell lung cancer; ET, etoposide; CRT, chemoradiotherapy; SRT, standard radiation therapy; HRT, hyperfractionated radiation therapy; MTD, maximum tolerated dose; PD-L1, programmed death-ligand 1; IHC, immunohistochemistry; ORR, objective response rate; DLT, dose-limiting toxicity; AEs, adverse events; CT, chemotherapy; CBT, chemobiotherapy; CRT, chemoradiotherapy; ORR, overall response rate; mOS, median overall survival; mPFS, median progression-free survival; DCR, disease control rate.

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durvalumab with or without tremelimumab after chemotherapy alone in ES-SCLC patients is currently active and not recruiting patients (NCT0304872). Its results are expected in 2020. The primary endpoints of the clinical trial include OS. The trial is designed as follows. In Arm A, durvalumab and tremelimumab are administered every 3 weeks for 12 weeks and carboplatin/cisplatin and etoposide are administered up to four cycles every 3 weeks. In Arm B, durvalumab is administered every 3 weeks for 12 weeks and carboplatin/cisplatin and etoposide are administered up to four cycles every 3 weeks. In Arm C, carboplatin/cisplatin and etoposide are administered up to six cycles every 3 weeks (Table 2).

Next generation of immune checkpoint targets

Immune checkpoint molecules are important regulators for maintaining immune homeostasis and preventing autoimmunity. These pathways consist of both stimulatory and inhibitory pathways, which are important for maintaining self-tolerance and regulating the type, magnitude, and duration of the immune response, as shown in Figure 1.

LAG3

Lymphocyte activation gene-3 (LAG3, CD223) is expressed by T cells and NK cells after MHC class II ligation. LAG3 and PD-1 are frequently co-expressed and upregulated on tumor-infiltrating lymphocytes, leading to immune exhaustion and tumor growth. LAG-3 blockade improves tumor immune responses and potentiates a variety of forms of immunotherapy to initiate different mechanisms of action that are mediated by impeding cell cycle progression. A combinational therapy of PD-1 with anti-LAG3 is an effective tool to treat many cancer types and produces a synergistic effect. However, the combined effects of other immune checkpoint inhibitory molecules and anti-LAG3 antibody are currently under investigation.

LAG525 is an immunoglobulin G4 humanized monoclonal antibody that binds LAG-3 with low nanomolar affinity, inhibiting the LAG-3 interaction with MHC class II and potentially restoring the activity of antitumor effector cells and enhancing anti-PD-1 antitumor activity. Anti-LAG3 monoclonal antibody LAG525 is being studied in a phase II clinical trial of patients with advanced solid and hematologic malignancies including SCLC (NCT03365791). Recruitment is currently underway. The primary endpoint is the clinical benefit rate at 24 weeks of PDR001+LAG525 and the PFS. PDR001 and LAG525 will be administered via i.v. infusion over 30 minutes once every 3 weeks. The results should be available in 2021 (Table 3).

OX40

OX40 (CD134) is a member of the TNF receptor superfamily that is highly expressed by activated CD4, CD8 T cells, and Tregs and less expressed by neutrophils and NK cells. OX40 is a secondary co-stimulatory immune checkpoint molecule that is expressed for 24 to 72 hours following its activation. Its ligand, OX40L, is also not expressed on resting APCs. OX40 and OX40L play a pivotal role in the activation, potentiation, proliferation, and survival of T cells and in modulating NK cell function. PF-04518600 (PF-8600) is an IgG2 humanized agonistic monoclonal antibody of OX40 that is being evaluated in the international JAVELIN Medley clinical trial (NCT02554812). The primary endpoint is the number of participants with dose-limiting toxicities in the first 8 weeks of treatment and those with an objective response. Currently, no data are unavailable on SCLC (Table 3).

4-1BB

4-1BB, an activation-induced co-stimulatory receptor, is an important regulator of immune responses that is expressed by T cells, NK cells, and APCs. Once expressed, it binds to its ligand (4-1BBL) and triggers the proliferation and activation of immune cells, particularly T and NK cells. Utomilumab (PF-05082566), a fully human monoclonal antibody that stimulates 4-1BB, is currently being studied in patients with SCLC and in combination with avelumab and PF-04518600 in a phase II clinical trial (JAVELIN Medley). The primary endpoint is the number of participants with dose-limiting toxicities in the first 8 weeks of treatment and those with an objective response. Utomilumab is administered to those who have progressed after at least one line of platinum-containing therapy, and one of three dose levels is to be used to optimize the combination with avelumab until disease progression. No preliminary results are available yet (Table 3).

Non-t cell-associated inhibitory molecules TGF-β

Transforming growth factor (TGF)-β is a cytokine that controls multiple cellular processes such as growth suppression of epithelial cells, alveolar epithelial cell differentiation, fibroblast activation, and extracellular matrix organization. The TGF-β pathway promotes cancer progression by
Table 3 Ongoing trials: next generation of immune checkpoint agents in SCLC

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<td>NCT03361228</td>
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<td>VII</td>
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Abbreviations: SCLC, small cell lung cancer; PFS, progression-free survival; ORR, objective response rate; DLTs, dose-limiting toxicities; TEAE, treatment-emergent adverse event; OR, objective response; CBR, clinical benefit rate; CR, complete response; PR, partial response; CRR, complete response rate.
concomitantly boosting tumor metastasis while inhibiting the host immune response. Although this pathway is able to control early-stage tumors by promoting cell cycle arrest and apoptosis, in advanced stages, it allows for tumor evasion by suppressing cytotoxic T cells and promotes cancer cell proliferation, invasion, and metastasis, a functional switch known as the TGF-β paradox. M7824 is a monoclonal antibody against PD-L1 linked to the extracellular domain of human TGF-β receptor 2 protein. Upon administration, the PD-L1 monoclonal antibody part of M7824 binds to PD-L1 while the TGF-β part binds to and neutralizes TGF-β receptor 2 protein, which prevents TGF-β- and PD-L1-mediated signaling and promotes CD8+ T cell and NK cell activation, ultimately inhibiting tumor growth. M7824 is being evaluated in a phase I/II clinical trial (NCT03554473) in patients with relapsed SCLC. The primary endpoints include efficacy (after 6 weeks for Arm B and 8 weeks for Arm C), PFS, duration of response, and OS. M7824 is administered at 1800 mg i.v. over 1 hour every 3 weeks on a 3-week cycle (Arm A/B) and at 1200 mg i.v. over 1 hour every 2 weeks on a 4-week cycle (Arm C). Topotecan is administered at 1 mg/m² i.v. over 30 minutes on days 1–5 on a 3-week cycle. Temozolomide is orally administered at 200 mg/m² on days 1–5 on a 4-week cycle. The results are expected in 2022 (Table 3).

IDO
Indoleamine 2,3-dioxygenase (IDO) is an intracellular enzyme that converts tryptophan to kynurenines. Kynurenines promote the differentiation and activity of Treg cells and decrease the amount and activity of CD8 T cells, which prompts an immunosuppressive situation that is compounded by the elevated amounts of PD-1/PD-L1 present in this milieu. Epacadostat is an oral agent that blocks the IDO pathway and is being investigated in a phase I/II clinical trial of advanced and metastatic malignancies including SCLC (NCT03277352). The primary endpoints include treatment-emergent adverse events, objective response rate, and the complete response rate. Treatment-emergent adverse events are screened through 60 days after the end of treatment up to 18 months for phase I. The objective response rate and complete response rate are to be assessed every 9 weeks for 12 months, then every 12 weeks up to 18 months for the phase II trial. Moreover, a non-randomized, open-label, phase I/II clinical trial of advanced solid tumors including SCLC (NCT03361228) is currently active and not recruiting patients. The primary endpoints include treatment-emergent adverse events and objective response rate, assessed up to approximately 12 months. No preliminary results are available yet (Table 3).

Conclusion
Immune checkpoint inhibitors are showing promise as therapeutic agents for SCLC and are being intensively studied. Thus, numerous trials at all stages and lines of treatment of SCLC are exploring the value of these agents both alone and in combination with chemotherapy or targeted agents. Overall, the expectations for immuno-oncology are high and the outcomes of these trials will hopefully reveal a variety of treatment options for SCLC patients. The discovery of new immune inhibitory and stimulatory pathways and rational combination strategies as discussed in this article is essential and will help to improve SCLC immunotherapy.

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Disclosure
The authors report no conflicts of interest in this work.

References


