Immune Checkpoint Inhibitor Therapy Achieved Complete Response for Drug-Sensitive EGFR/ALK Mutation-Negative Metastatic Pulmonary Large-Cell Neuroendocrine Carcinoma with High Tumor Mutation Burden: A Case Report

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Abstract: Large-cell neuroendocrine lung carcinoma (LCNELC) is classified into lung neuroendocrine tumors according to WHO 2015 classification guidelines and represents approximately 3% of all lung cancer. Because of the rarity of LCNELC, there is a lack of prospective studies guiding treatment. Here, we report a case of a patient with pT2aN2M0 stage IIIA LCNELC (drug-sensitive EGFR/ALK mutation-negative, PD-L1-negative but tumor mutation burden (TMB) high), who progressed rapidly after surgery but achieved a complete response to subsequent immune checkpoint inhibitor (ICI) therapy. The concentration of circulating tumor DNA (ctDNA) following the treatment course strongly reflects the response to ICI therapy. This report highlights the efficacy of ICI treatment in metastatic LCNELC patients with a high TMB and suggests that ctDNA analysis in detecting molecular residual disease may facilitate the personalization of ICI therapy.

Keywords: metastatic large-cell neuroendocrine lung carcinoma, adrenal gland metastasis, next generation sequencing, immune checkpoint inhibitor, circular tumor DNA

Introduction

Large-cell neuroendocrine lung carcinoma (LCNELC) is a poorly differentiated tumor that exhibits large cells with neuroendocrine features and a low nuclear-to-cytoplasmic ratio, representing approximately 3% of all lung cancer.1 Because of the rarity of LCNELC, there is a lack of prospective studies or strong evidence to guide treatment. The primary treatment strategy for early-stage LCNELC patients is complete resection with mediastinal lymph node dissection, but the disease often rapidly recurs.2–4 Therefore, efficient adjuvant therapy after surgery is necessary for LCNELC patients.5 However, a multi-center retrospective study from China showed that patients with LCNELC had a progression-free survival (PFS) of only 11.5 and 7.2 months, respectively, when receiving small-cell lung cancer (SCLC)-based regimens and non-small-cell lung cancer (NSCLC)-based regimens.6 The inhibition of the antigen-specific T-cell responses by the immune checkpoint leads to tumor immune escape,7 which can be corrected by immune checkpoint inhibitors (ICIs) though reversing the tumor-derived inhibitory immune reactions or activating anti-tumor immune...
activities. Here, we report a metastatic LCNELC patient with a high tumor mutation burden (TMB) level who benefited from ICI treatment and achieved a PFS of 20 months.

Case Report
We report a 54-year-old man with a 40 pack-year history of smoking, who underwent a routine examination in July 2018. The patient gave written informed consent for publication of his case details and images. The publication of the case details was approved by the ethics committee of The First Hospital of China Medical University (AF-SOP-07-1). Computed tomography (CT) of the chest revealed a lung lesion measuring 4 cm in the right upper lobe with No.4R lymph node enlargement (Figure 1A and B). No distant metastases were found through brain magnetic resonance imaging (MRI), full-body bone scan, and ultrasonography of the abdomen and adrenal glands. Because of bullae in the lung around the tumor and the risk of pneumothorax after biopsy, CT-guided needle biopsy was rejected by the patient. A lobectomy and lymph node dissection were performed in August 2018, and pT2aN2M0 stage IIIA LCNELC with No.4R lymph node metastasis was confirmed via pathological diagnosis. Immunohistochemical analysis showed positive staining for CK, TTF-1, synaptophysin, PAS, chromogranin A and Ki-67 (80%), focal positive staining for CK7, CD56 and CD117, but negative staining for CK5/6, P40, P63, CD45 (LCA) and PSA-AB, consistent with a LCNELC origin (Figure 2). Adjuvant chemotherapy was rejected by the patient.

To seek a more effective treatment strategy, tumor and plasma biopsies were subjected to genetic testing using targeted next generation sequencing (NGS) for 425 cancer-relevant genes (Gene seqPrime) and numerous tumor-specific mutations (Table 1) were observed with a high TMB at 25.8 mutations per MB (Method in Supplement 1). However, no drug-sensitive EGFR mutation or ALK rearrangement was identified.

After surgery (2 months), chest and abdomen CT with contrast enhancement was conducted as the baseline before adjuvant therapy. The scans revealed a 2-cm-sized metastatic tumor in the left adrenal gland, which indicated progressive disease (Figure 3A). Although immunohistochemical staining of the primary tumor for the programmed death-ligand 1 (PD-L1) was negative (Figure 4), ICI therapy was recommended to the patient because of the high level of TMB and negative mutation of EGFR/ALK. Then the patient began to receive nivolumab treatment (240 mg) in October 2018.

The efficiency of the treatment was evaluated after every 4 cycles of treatment through abdomen CT and neuron-specific enolase (NSE) detection. During 15 cycles of treatment, the patient achieved a complete response (CR), with significant regression of the metastatic tumor in the left adrenal gland (Figure 3B–E). Before the 16th cycle of treatment, the patient suffered from cough with mild dyspnea and a chest CT showed interlobular septal thickening and subpleural ground-glass opacities (Figure 5A). Nivolumab therapy was discontinued for 3 months but the symptoms and conditions revealed by the CT images became worse (Figure 5B). By treating the patient with acetylcysteine and corticosteroids for 3 months, the symptoms disappeared and the CT imaging results recovered (Figure 5C and D). By 4, 8 and 12 months after drug withdrawal, no progression occurred (Figure 3F–H) and the level of NSE remained in the normal range (Figure 6A). In addition, the concentration of

Figure 1 The CT scan of the patients before surgery. (A) The 4 cm tumor on the right upper lobe (Red arrow); (B) Metastatic right No.4 lymph node (Red arrow).
circulating tumor DNA (ctDNA) from three sequential plasma samples collected before, and two and four cycles after nivolumab treatment showed a continued decrease during the treatment course, which indicated a durable response to nivolumab (Figure 6B). During 20 months of post-treatment follow-up (until May 2020), no recurrence or metastasis was observed in the patient.

**Discussion**

According to the Surveillance, Epidemiology and End Results (SEER) database, LCNELC is a rare disease that is more commonly detected in elderly and male patients, and in the upper lobe. LCNELC shares similarities with SCLC on the basis of clinical and pathological features, while LCNELC patients who have previously undergone complete resection and received adjuvant chemotherapy with the standard combination regimen of SCLC (cisplatin plus etoposide) show prolonged survival. However, therapeutic approaches for metastatic and unresectable tumors remain to be further evaluated. Recently, the efficacy of ICIs has been confirmed in various treatment approaches for neuroendocrine tumors. A retrospective cohort study in France showed that patients with LCNELC had a significant response to immune checkpoint inhibitors and achieved a median PFS of 57 weeks. In the case we report here, the patient exhibited a CR and achieved a PFS of 20 months (until May 2020) upon ICI treatment.

**Figure 2** Immunohistochemical staining. (A) Hematoxylin-eosin staining of the tumor tissue (40X and 400X); (B) Immunohistochemical staining for Synaptophysin (Positive, 400X); (C) Immunohistochemical staining for Chromogranin A (Positive, 400X); (D) Immunohistochemical staining for Ki-67 (80% Positive, 400X).

### Table 1 Mutations Detected by NGS

| Gene   | Mutation             | Gene   | Mutation            |
|--------|----------------------|--------|---------------------|
| AKT2   | p.E342K(c.G1024A)    | HGF    | p.D275H(c.G823C)    |
| ARID1A | p.G171I(c.G1537A)    | LRP1B  | p.V4161F(c.G12481T) |
| ARID2  | p.D396Y(c.G1186T)    | NX2-1  | p.A2975T(c.G899T)   |
| BUB1B  | p.G831V(c.G2492T)    | PARP2  | p.Q177H(c.G531C)    |
| CDRK2A | p.D108Y(c.G322T)     | PDGFRB | p.D691Y(c.G2071T)   |
| DDR2   | p.G517Y(c.G1550T)    | PIK3C3 | p.E646H(c.G1938G)   |
| DUSP2  | p.E468fs(c.G1396delG)| QKI    | p.A219T(c.C665A)    |
| EGFR   | p.E858Q(c.G2602C)    | RFI    | p.S230X(c.G689A)    |
| EPHA3  | p.A514D(c.C1541A)    | ROS1   | p.P1053L(c.C3158T)  |
| EXT2   | p.D540H(c.G1618C)    | TAP2   | p.P6365fs(c.C1906T)|
| FANCA  | p.R7566(c.G2267T)    | TP53   | p.R280K(c.G839A)    |
| FGFR1  | p.H1164N(c.C490A)    | YAP1   | p.D93N(c.G277A)     |
highly associated with pre-existing interstitial lung disease.\textsuperscript{16,17} ICI-ILD often occurs earlier in lung cancer, with a median of 2.1 months after ICI treatments,\textsuperscript{1} and presents as several radiologic patterns.\textsuperscript{18} A total of 70–80\% of ICI-ILD cases can be controlled through the prompt initiation of a high dose of steroids.\textsuperscript{19} In our case, the male patient had a long history of smoking and his chest CT showed severe pre-existing interstitial lung disease, which might lead to the occurrence of ILD after 15 cycles of nivolumab treatment.

High TMB is regarded as a predictive biomarker of response to ICIs.\textsuperscript{20,21} It has been reported that TMB in LCNELC is commonly higher compared with other NSCLC subtypes,\textsuperscript{22} which indicates that ICIs might be an option for LCNELC. Wang et al reported a LCNELC patient with a high TMB who benefited from ICI treatment.\textsuperscript{23} In our case, the patient was PD-L1 negative but had a high TMB, at 25.6 mutations per Mb, and achieved a CR after ICI treatment, suggesting that immunotherapy is effective in this class of disease.

The level of ctDNA in molecular residual disease detection during treatment strongly predicts recurrence for multiple tumor types.\textsuperscript{24} In our case, the concentration of ctDNA continued to decrease after two cycles.
of ICI treatment and was no longer detectable after four cycles. Follow-up imaging demonstrated consistent results, with no evidence of progressive disease 20 months after starting ICI treatment. Our findings suggest that ctDNA analysis could potentially guide the decision to distinguish responders at an early stage during ICI therapy in LCNELC patients.

Conclusion
In summary, we reported an advanced LCNELC patient with a high TMB level who achieved a CR on nivolumab treatment with a significant decrease of ctDNA concentration during treatment and achieved a PFS of 20 months. This report highlights the efficacy of ICI treatment in metastatic LCNELC patients with a high TMB and suggests that ctDNA analysis may facilitate the personalization of ICI therapy.

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

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