Sustained Oligoclonal T Cell Expansion Correlates with Durable Response to Immune Checkpoint Blockade in Lung Cancer

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Abstract

Purpose: Antibodies that target immune checkpoint molecules have demonstrated significant and durable clinical benefit through re-activation and proliferation of pre-existing tumor-infiltrating CD8+ T cells in a broad range of tumor types including lung cancer. Detailed characterization of T cell dynamics at clonal levels using next-generation T cell receptor (TCR) sequencing in large cohorts of lung cancer patients is yet reported.

Methods: We performed TCR sequencing of peripheral blood samples (and some tumors) obtained from 27 lung cancer patients undergoing single/combined immune checkpoint blockade therapies.

Results: In one responder, we found expansion of a single T cell clone (approximately 20% of the all TCR reads) in a metastatic subcutaneous lesion that showed pathologic complete response on day 17 of treatment. The same TCR CDR3 sequence was detected in 6.1% of TCR reads in the peripheral blood prior to treatment initiation and the expansion remained persistent in peripheral blood at 21.0% at week 10 and 24.3% at week 48. In other patients who showed durable response, we also found persistent oligoclonal T cell expansion in their peripheral blood which was not observed in non-responders even while they remained on therapy.

Conclusion: We found sustained expansion of oligoclonal T cell clones in lung cancer patients who had durable response to immune checkpoint blockade. This suggests possible use of longitudinal TCR sequencing to assist clinical decision-making, assess synergism with other agents, and most importantly facilitate rational development of alternative treatment strategies for non-responders.

Keywords: Immune checkpoint blockade; Immunotherapy; Lung cancer; Next-generation sequencing; T cell receptor

Introduction

Following decades of skepticism, the prominent roles of an immune system in tumor control are evidently proven through the unprecedented durable response shown by immune checkpoint inhibitors in many tumor types. Characterization of the complex interactions among tumor cells, antigen-presenting cells, and effector T cells at molecular levels revealed key pathways in an immune signaling that mediate tumor escape from host immune surveillance. First was the discovery of a cytotoxic T lymphocyte antigen-4 (CTLA-4) molecule which was shown to down-modulate T cell receptor (TCR) signaling following stimulation of naïve T cells [1,2]. Activation of anti-tumor effects through inhibition of CTLA-4 and possible mechanisms of action were later demonstrated in mouse models followed by clinical development [3-5]. Programmed death-1 (PD-1) is expressed on T cells upon their activation [6,7] and plays a major role in maintenance of tolerance by interaction with its ligand PD-L1 which suppresses immune-mediated tissue damage [8]. Clinical trials of antibodies targeting the PD-1/PD-L1 pathway demonstrated improvement in median overall survival when compared to conventional chemotherapies [9,10] and led to approvals of nivolumab, pembrolizumab and atezolizumab for non-small cell lung cancer (NSCLC). However, the fact that the majority of patients did not have any clinical benefit from these agents necessitates the discovery of an accurate biomarker to guide patient selection. Both PD-L1 expression and mutational burden were suggested to be correlated with the response to immune checkpoint blockades, but the results are still controversial and their clinical use is very limited [9,11,12]. In the trial of first-line pembrolizumab for NSCLC, the majority of patients (55.2%) did not respond in spite of high expression of PD-L1 (>50%) or higher mutational burden [12,13]. These immune checkpoint blockades do not act directly on cancer cells, but eradicate cancer cells through the activation of cytotoxic T cells that target cancer cells. Hence, comprehensive longitudinal assessment of anti-tumor T cells should provide better insight into the anti-tumor effect of these agents and also may lead to the identification of factors essential for clinical benefit. Immunohistochemistry of serial biopsies obtained from responders has shown increased infiltration of CD8+ T cells in the tumor microenvironment [14,15]. The assessment of these processes at clonal levels of T cells has recently been achieved by advances in high-throughput sequencing technologies [16,17], which have enabled us to identify millions of individual TCR sequences and provide the opportunity to track each of them longitudinally. Previous analyses of samples at two time-points (before and after the treatment) in a small cohort of patients revealed correlation between clonal expansion of certain T cells and clinical responses to immune checkpoint blockades [18-20]. Such findings are consistent with other reports which suggested

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tumor eradication by immune checkpoint blockade through T cell clones targeting tumor-specific antigens [21,22]. Therefore, a detailed study of T cell clonal dynamics in patients with the assessment of both immediate expansion and long-term persistence in responders versus non-responders to immunotherapy could provide a valuable insight into the mechanisms that mediate tumor eradication and durable response by these agents.

Materials and Methods

Patients

We enrolled a total of 27 patients with lung cancer, who received single agent anti-PD-1 antibodies or a combined anti-PD-1 and anti-CTLA-4 antibodies, at University of Chicago Hospitals under the institutional review board (IRB) approval (#9571). These include patients with different lung cancer subtypes (small cell and non-small cell) and other thoracic malignancies (thymic and large cell neuroendocrine tumors). Detailed patient information is summarized in Table 1. We analyzed 2 or more serial peripheral blood samples obtained from each patient. Blood samples (10-20 mL) were collected in BD Vacutainer CPT Cell Preparation tubes (BD Biosciences, San Jose, CA, USA) and peripheral blood mononuclear cells (PBMCs) were separated according to manufacturer’s instructions. In patient A16.003, we also performed analyses on excess tumor samples obtained during routine clinical care as approved by IRB and with the consent of the patient. For patient A16.003 and A16.015, we separated CD8+ and CD8− cell fraction using Dynabeads CD8 Positive Isolation Kit (Thermo Fisher Scientific, Carlsbad, CA).

TCR repertoire analysis

Total RNA was extracted from tumor/PBMC samples using RNeasy mini kit (Qiagen, Valencia, CA, USA) and TCR sequencing was performed as previously described [17,19,23]. Briefly, we synthesized cDNA with 5’ rapid amplification of end (5’-RACE) adapter using SMART cDNA library kit (Clontech Laboratories, Mountain View, CA, USA). We then amplified the TCR cDNA with reverse primers specific for the constant regions of TCRα or TCRβ, and a forward primer for the SMART adapter. After adding Illumina sequence adapter with barcode sequences using the Nextera XT Index kit (Illumina, San Diego, CA, USA), we performed deep sequencing by 300-bp paired-end reads on the MiSeq platform (Illumina). We used Tcrip software for the TCR repertoire analysis as described previously [17].

Results

Patient characteristics

We analyzed TCR repertoire of 27 patients from whom we could obtain 2 or more serial samples (Table 1). All patients had stage IV diseases consisting of 16 lung adenocarcinomas, 6 squamous cell lung cancers, 2 small-cell lung cancers and 3 other thoracic malignancies. We obtained samples before immunotherapies in 13 (48.1%) of the 27 patients and analyzed an average of 3.3 time-points per patient. In this selected patient population, 44.4% were responders (R; 12/27) and 55.6% were non-responders (NR; 15/27), according to the RECIST criteria [24]. Among the responders, two patients (A16.003 and A16.010) did not previously receive any chemotherapy because of concerns about their capacity to tolerate therapy. The average number of previous lines of therapy was not different between the responder and non-responder groups (P=0.25; 1.33 vs 1.73). The distribution of lung cancer subtypes was similar between the two groups. The average age was higher among the non-responders (66.3 years) than responders (59.5 years), but not significantly different (P=0.15).

Oligoclonal T cell expansion in a tumor and peripheral blood in one responder

From a patient A16.003 with metastatic squamous cell lung cancer who had previously received palliative radiation therapy to his chest lesions (Figure 1A), we obtained biopsy of a subcutaneous lesion around the right antecubital fossa on day 17 after two cycles of anti-PD1 therapy (Figure 1B), by which time there was pathological complete response. We analyzed TCR repertoire by using our next-generation TCR sequencing method and found possible oligoclonal T-cell expansion that was reflected by a single dominant CDR3 sequence of TCRα and TCRβ in their sequence reads (Figure 1C). The CT scan at week 10 after receiving 4 cycles of therapy revealed significant tumor shrinkage (Figure 1D). We also analyzed TCR repertoire of peripheral blood in this patient at multiple time-points. Interestingly, even before the initiation of therapy, the peripheral blood sample of this patient possessed the TCR clonotype at frequency of 6.1% which same as was observed at the site of early pathological complete response. This specific clonotype expanded to frequency of 21.0% in week 10 and remained at the frequency of 24.3% at week 48 of the therapy (Figure 1E). We confirmed that the specific T cell clone was CD8+ T cells. This suggest that tumor eradication is mediated by reactivation and expansion of pre-existing tumor infiltrating T cell clone.

Table 1: Baseline patient characteristics.

<table>
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<th>Sample</th>
<th>Gender</th>
<th>Age</th>
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<th>Stage</th>
<th>Immunotherapy</th>
<th>Response</th>
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NSCLC-NOS: Non-small cell lung carcinoma-not otherwise specified; R: Responder; NR: Non-Responder


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Sustained oligoclonal T cell expansion in peripheral blood correlates with response

We further analyzed TCR repertoire in peripheral blood of the patients before and after treatment initiation (Figure 2). In two responders, A16.010 (Figures 2A and 2B) and A16.057 (Figures 2C and 2D), for whom we had samples before treatment initiation, we observed oligoclonal expansion of few dominant T cell clones after 1 cycle of the treatment in A16.010 and after 4 cycles of the treatment in A16.057. In both A16.010 and A16.057, 1-2 dominant T cell clones detected at early time-points were sustained at week 10 of the therapy. In the other patient A16.008 (Figures 2E and 2F), although we did not have a baseline analysis, there was a single dominant T cell clone that remained at the high frequency over 8 months of monitoring with a
Absence of T cell expansion in peripheral blood correlates with resistance to immune checkpoint blockade. (A, B) Patient A16.023. Her chest wall mass continued to progress even as she remained on anti-PD1 therapy (anti-CD137 antibody was added after 4 cycles of anti-PD1 therapy as part of clinical trial) (A). TCR repertoire analysis showed no evidence of T cell clonal expansion in her peripheral blood at week 2 and week 10 (B). (C, D) Patient A16.043. He did not respond to index immunotherapy as revealed on week 23 CT scan while still on therapy, although he had previously been received anti-PD1 therapy 3 years earlier as part of clinical trial during which he had 'stable disease' (C). TCR repertoire analysis showed no evidence of clonal expansion in his peripheral blood at week 12 (D). (E, F) Patient A16.022. There is evidence of progressive with increase in size of right lower lobe lesion (left lower lobe opacification persisted) (E). T cell clonal expansion was transient as most were back to baseline at week 12 of treatment (F). (G, H) Patient A16.015. Imaging of patient A16.015 revealed progressive disease at week 10 (G), and TCR repertoire analysis showed one T cell clonal was expanded at week 10; however, this clone was CD8+ (H).

Figure 3: Absence of T cell expansion in peripheral blood correlates with resistance to immune checkpoint blockade. (A, B) Patient A16.023. Her chest wall mass continued to progress even as she remained on anti-PD1 therapy (anti-CD137 antibody was added after 4 cycles of anti-PD1 therapy as part of clinical trial) (A). TCR repertoire analysis showed no evidence of T cell clonal expansion in her peripheral blood at week 2 and week 10 (B). (C, D) Patient A16.043. He did not respond to index immunotherapy as revealed on week 23 CT scan while still on therapy, although he had previously been received anti-PD1 therapy 3 years earlier as part of clinical trial during which he had 'stable disease' (C). TCR repertoire analysis showed no evidence of clonal expansion in his peripheral blood at week 12 (D). (E, F) Patient A16.022. There is evidence of progressive with increase in size of right lower lobe lesion (left lower lobe opacification persisted) (E). T cell clonal expansion was transient as most were back to baseline at week 12 of treatment (F). (G, H) Patient A16.015. Imaging of patient A16.015 revealed progressive disease at week 10 (G), and TCR repertoire analysis showed one T cell clonal was expanded at week 10; however, this clone was CD8+ (H).

stable disease condition seen on imaging obtained at week 41. Other patients with durable response showed similar pattern of sustained dominant T cell clones during several months of monitoring period (Supplementary Figures 1A-1H). In contrast, non-responders did not show any dominant T cell clonal expansion or no persistence of dominant T cell clones in those we did not have baseline samples on (Figures 3A-3H) (Supplementary Figures 2A-2L). Although patient A16.015 who showed oligoclonal T cell expansion at week 10 when his tumor did not show any response to anti-PD-1 therapy, we confirmed this T cell clone were not CD8+ T cells (Figures 3G and 3H). These data further suggest that early T cell clonal expansion in peripheral blood may correlate with durable clinical response.

Discussion

The elucidation of mechanisms through which tumors evade surveillance by host immune system through immune checkpoints has ushered in a new era in oncology. A subset of patients with advanced metastatic melanoma treated with agents blocking such immune checkpoints has lived beyond a decade [25]. Such significant clinical benefit has led to approvals of these agents for many tumor types. However, the fact that only a fraction of the patient population derives clinical benefits has necessitated the development of biomarkers that will guide patient selection, detect response to therapy, and determination of optimal duration of therapy. Detailed characterization of mechanisms of tumor eradication in responders at cellular levels may provide blueprint for alternative treatment strategies for non-responders. Studies in mouse models showed that these agents enhance functional activity and proliferation capability of cancer specific cytotoxic T cells [26]. The expanded T cells were demonstrated to target neoantigens generated by somatic mutations and immunization with a mutant peptide(s) was shown to cause tumor rejection similar to immune checkpoint blockade [21]. These findings were further supported by a report showing that a responder to a CTLA-4 blockade had T cell clonal expansion targeting a neoantigen [22]. Thus, it is likely that tumor eradication by these agents is through reactivation and proliferation of pre-existing tumor infiltrating cancer-specific cytotoxic T cells. This is concordant with the results that ex-vivo expansion and adoptive transfer of mutant-specific T cell clones achieved durable clinical response in different tumor types [27,28]. Hence, longitudinal comprehensive analyses of T cell dynamics both at the tumor site(s) and in peripheral blood may elucidate the footprints of tumor eradicating T cell dynamics in responders. Recent advances in the next-generation sequencing have provided us tools to measure dynamic changes of immune cells including individual T cell clones within tumors and peripheral blood samples [27]. In our patient cohort, we found a patient who had a single dominant T cell clone with the TCRαβ frequencies of around 20% (Figure 1C) at a metastatic site with pathological complete response at day 17 of the therapy. The same T cell clone was found in peripheral blood at the frequency of 6.1% before the treatment and was expanded to 21.0% at week 10. This frequency is similar to that reported in a melanoma patient who showed complete radiological response [20], thus monitoring T cell clonal dynamics using this comprehensive approach (i.e. analyses of entire repertoire) may reflect response to therapy more accurately. Our data also suggest that oligoclonal T cell clonal expansion after one or a few cycles of therapy may be an early sign of favorable clinical response. Although we need a larger number of samples at multiple fixed time-points in comparison to the baseline pretreatment sample and further confirmation that those expanded T cells are reactive to cancer cells, monitoring T cell repertoire in peripheral blood could expedite clinical decision making in neoadjuvant and oligometastatic disease settings. Patients with evidence of early oligoclonal T cell expansion may remain on therapy while those without clonal T cell expansion may proceed to surgery or start on alternative therapies. Similar to previous
reports in mouse models and in human adoptive T cell transfer therapies [21,27,28], we found that 1-2 dominant T cell clones may be responsible for tumor eradication in responders and their persistence at high frequencies correlate with durable response (Figures 1a and 2). Even in the patients for whom we did not have baseline pre-treatment samples but who had durable response, we observed the persistence of 1-2 dominant T cell clones in their peripheral blood for many months (Supplementary Figure 1). This further supports previous observations that only a subset of patients that had adoptive T cell therapy had durable response and in them (unlike non-responders) the re-infused T cell clones following ex-vivo expansion remained at high frequencies for several months afterwards [27,28]. So also a recent report demonstrated that non-responders to immunotherapy actually had T cell clones targeting neoantigens, but yet did not derive clinical benefit like as a result of the inability of these T cell clones to proliferate [29]. These together suggest that in addition to having cancer-specific tumor infiltrating T cells, it is also critically important for these T cells to have self-renewal and sustained expansion capacities. The response rate in this study cohort did not reflect correctly in an entire clinic since we included patients whom we already knew to have durable response.

**Conclusion and Limitations**

The limitations of our study include our inability to identify antigens targeted by the expanded T cell clones due to unavailability of tissue samples for tumor exome sequencing. We cannot exclude a possibility that some of the T cell clonal expansions may be caused by other factors (e.g. infections). However, the pattern of T cell clonal expansion by infection would be typically rapid followed by contraction once the infection is controlled. In addition, we were unable to apply any statistical analyses to evaluate significant differences between responders and non-responders because i) we only had three responders of whom we had baseline pre-treatment samples, ii) the time-points of sample collection varied among patients for logistical reasons, and then iii) we did not observe any correlation between tumor burden and clonal frequencies. We also need to consider immune suppression mechanisms in cancer cells using a large cohort set of patients. To the best of our knowledge, this is the largest report of longitudinal TCR sequencing of peripheral blood in patients treated with immunotherapy. It is also the first to include multiple time-points during the course of treatment from the first treatment to as far as 13 months (although we had no baseline samples in some of these cases). It is notable that early and sustained oligoclonal T cell expansion may be correlate with better durable response to immune checkpoint blockades and may provide deeper insights into mechanisms of tumor eradication by these agents.

**Conflict of Interest**

Y.N. is a stockholder and an advisor of OncoTherapy Science Inc. J.P. is a scientific adviser of OncoTherapy Science Inc. Other authors have no conflict of interest concerning this study.

**Acknowledgments**

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