Dynamics of Tumor and Immune Responses during Immune Checkpoint Blockade in Non–Small Cell Lung Cancer

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Abstract

Despite the initial successes of immunotherapy, there is an urgent need for molecular assays that identify patients more likely to respond. Here, we report that ultra-sensitive measures of circulating tumor DNA (ctDNA) and T-cell expansion can be used to assess responses to immune checkpoint blockade in metastatic lung cancer patients (N = 24). Patients with clinical response to therapy had a complete reduction in ctDNA levels after initiation of therapy, whereas nonresponders had no significant changes or an increase in ctDNA levels. Patients with initial response followed by acquired resistance to therapy had an initial drop followed by recrudescence in ctDNA levels. Patients without a molecular response had shorter progression-free and overall survival compared with molecular responders [5.2 vs. 14.5 and 8.4 vs. 18.7 months; HR 5.36; 95% confidence interval (CI), 1.57–18.35; P = 0.02, respectively], which was detected on average 8.7 weeks earlier and was more predictive of clinical benefit than CT imaging. Expansion of T cells, measured through increases of T-cell receptor productive frequencies, mirrored ctDNA reduction in response to therapy. We validated this approach in an independent cohort of patients with early-stage non–small cell lung cancer (N = 14), where the therapeutic effect was measured by pathologic assessment of residual tumor after anti-PD1 therapy. Consistent with our initial findings, early ctDNA dynamics predicted pathologic response to immune checkpoint blockade. These analyses provide an approach for rapid determination of therapeutic outcomes for patients treated with immune checkpoint inhibitors and have important implications for the development of personalized immune targeted strategies.

Significance: Rapid and sensitive detection of circulating tumor DNA dynamic changes and T-cell expansion can be used to guide immune targeted therapy for patients with lung cancer. See related commentary by Zou and Meyerson, p. 1038

Introduction

Despite the durable clinical benefit observed with immune checkpoint inhibitors for patients with non–small cell lung cancer (NSCLC), the majority of patients are either refractory or eventually develop acquired resistance after an initial response (1). Similar to the targeted therapy paradigm, success...
of immuno-oncology seems to depend on choosing patient populations most likely to benefit. The plasticity of the immune system under immunotherapy has weakened single biomarker-driven approaches (2) and currently used predictive biomarkers have been unable to accurately identify the subset of patients who benefit from these therapies.

We hypothesized that noninvasive molecular analyses that evaluate tumor-derived cell free circulating tumor DNA (ctDNA) and tumor-extrinsic T-cell receptor (TCR) repertoire parameters may be useful for rapidly determining which patients would ultimately benefit from immune checkpoint blockade. Such approaches may be of particular importance for immune-targeted agents as the therapeutic responses have been challenging to evaluate using radiographic imaging due to tumor immune infiltration (3). Conventional response criteria such as the RECIST do not consistently capture the unique patterns and timing of antitumor immune responses (4, 5).

The temporal relationship between detection of ctDNA and emergence of recurrent or progressive disease has been shown in patients with early-stage NSCLC (6, 7) and as we show in our companion study in patients with advanced NSCLC receiving targeted therapies (8). During treatment with immunotherapy, our group has shown that ctDNA may be predictive of outcome in melanoma patients treated with CTLA-4 blockade (9). ctDNA changes have been associated with therapeutic outcome during immune checkpoint blockade in NSCLC (10–12), however these analyses have been limited by the low sensitivity of the approaches, permitting analyses in approximately half of the cases analyzed. Even less is known about the dynamics of the peripheral T-cell repertoire during immune checkpoint blockade in NSCLC (13) and how these changes relate to ctDNA levels and tumor response.

To overcome these issues and to allow ultrasensitive evaluation of ctDNA during therapy, we have developed targeted error-correction sequencing (TEC-Seq), a custom capture and sequencing approach that permits sensitive and specific detection of low abundance sequence alterations using next-generation sequencing (14). We have also developed new methods of evaluating TCR clonal expansion in the tumor microenvironment during immune checkpoint blockade (13). Here, we use these approaches to investigate whether ctDNA and TCR dynamics are reflective of therapeutic outcome for patients with NSCLC treated with immune checkpoint blockade.

**Patients and Methods**

**Patient characteristics**

Our study group consisted of 24 metastatic patients with NSCLC treated with immune checkpoint blockade as a standard of care (n = 19) or in the setting of a clinical trial (n = 5) between October 2014 and August 2016 at Johns Hopkins Sidney Kimmel Cancer Center. In parallel, we evaluated 14 patients with stage 1 to IIIA surgically resectable NSCLC that received anti-PD1 therapy in the setting of a neoadjuvant nivolumab clinical trial (15). The studies were conducted in accordance with the Declaration of Helsinki, were approved by the Institutional Review Board (IRB), and patients provided written informed consent for sample acquisition for research purposes. Clinical characteristics for all patients are summarized in Supplementary Table S1.

**Treatment and assessment of therapeutic response**

Therapeutic responses were evaluated by the RECIST version 1.1 (16). Baseline disease burden was determined by the sum of the longest diameters of target lesions as determined by RECIST 1.1 criteria. After baseline imaging, radiographic evaluation was performed at 5- to 10-week intervals or as clinically indicated for the metastatic cohort and 7 days prior to surgery for the early-stage cohort. The timing of radiologic assessments typically followed the early timepoints of blood sample collection. Although this approach may be subject to lead-time bias, we sought to mirror the imaging schedule used in clinical practice. Furthermore, in contrast to chemotherapy or targeted therapy, where therapeutic response may be accurately evaluated by imaging early after treatment initiation, the unique nature and timing of response to immune checkpoint blockade mandates response assessments at later timepoints including confirmation of the radiologic response. Of the 19 metastatic patients with NSCLC with detectable ctDNA, one achieved complete response (CGLU111), three patients achieved partial response (CGLU1135, CGLU1337, and CGLU1347), and 12 achieved SD (CGLU1115, CGLU1117, CGLU1159, CGLU160, CGLU162, CGLU168, CGLU203, CGLU211, CGLU212, CGLU340, CGLU351, and CGLU357) as best overall response. Three patients (CGLU1121, CGLU1243, and CGLU1348) experienced disease progression. Of the three patients with partial response, two eventually developed molecular resistance. In the neoadjuvant cohort, a repeat chest CT ≤7 days prior to surgery revealed stable disease for all patients with detectable ctDNA at baseline.

**Blood sample collection**

For all patients, at least two serial blood samples (range 2–8) were collected over the course of treatment for isolation of plasma and extraction of cell-free DNA for genomic analyses. We analyzed a total of 105 serial plasma samples that were obtained prior to anti-PD1, at 4 to 8 weeks and additional time points during therapy for all metastatic patients with NSCLC except for CGLU1135 and CGLU1161. For these two patients, blood from early timepoints was not available and blood samples from the time of radiographic response and the time of acquired resistance were analyzed. A detailed description of the time points analyzed is shown in Supplementary Tables S2 and S3. Baseline tumors were analyzed by whole exome sequencing or targeted next-generation sequencing for patients in the metastatic cohort, with the exception of CGLU1168, for which a tumor specimen from the time of resistance to immune checkpoint blockade was used. For the early-stage cohort, tumor samples prior to therapy initiation or at the time of resection in the cases where baseline tumor was not available, were analyzed by whole exome sequencing (15).
Sample preparation and next-generation sequencing of cfDNA

Whole blood was collected in K2 EDTA tubes; plasma and cellular components were separated by centrifugation at 800 × g for 10 minutes at 4°C. Plasma was centrifuged a second time at 18,000 × g at room temperature to remove any remaining cellular debris and stored at –80°C until the time of DNA extraction. DNA was isolated from plasma using the Qiagen Circulating Nucleic Acids Kit (Qiagen GmbH). TEC-Seq next-generation sequencing (CEL-Seq) DNA libraries were prepared from 12 to 125 ng of cfDNA. Genomic libraries were prepared as previously described and targeted capture was performed using the Agilent SureSelect reagents and a custom set of hybridization probes targeting 58 genes, described in Supplementary Table S4 (14). TEC-Seq libraries were sequenced using 100 bp paired end runs on the Illumina HiSeq 2500 (Illumina). The analytical performance and validation including sensitivity and specificity and limits of detection of our cfDNA platform have been recently reported (14).

Primary processing of cfDNA next-generation sequencing data and identification of putative somatic mutations

Primary processing of next-generation sequencing data for cfDNA samples was performed as previously described (14) using Illumina CASAVA software (v1.8), including demultiplexing and masking of dual index adapter sequences. Sequence reads were aligned against the human genome reference (hg19) using Novoalign with additional realignment of select regions using the Needleman–Wunsch method (17). Next, candidate somatic mutations, consisting of point mutations, small insertions, and deletions were identified using VariantDx (17) across the targeted regions of interest. VariantDx examined sequence alignments of cfDNA plasma samples while applying filters to exclude alignment and sequencing artifacts as previously described (14). Specifically, an alignment filter was applied to exclude quality failed reads, unpaired reads, and poorly mapped reads in the plasma. A base quality filter was applied to limit inclusion of bases with reported Phred quality score >30. Criteria for calling alterations in cfDNA have been previously described (14). TEC-Seq characteristics are shown in Supplementary Table S5.

Identification of tumor-derived cfDNA

Genomic alterations in cfDNA were cross-referenced against each patient’s tumor-specific genomic alterations to identify bona fide tumor-specific cfDNA variants. Variants identified in cfDNA as previously described (14) and in the matching tumor with a MAF of ≥2% were considered tumor specific. We focused on somatic variants that were identified both in the tumor sample and in cfDNA for each patient to exclude variants related to clonal hematopoiesis (Supplementary Table S6). Our dataset is deposited in the database of Genotypes and Phenotypes (dbGaP; study ID 32485).

Clonality estimates of cfDNA variants

To assess the cellular prevalence of plasma mutations in their corresponding tumors, tumor samples of each case were analyzed as follows. The density of reads mapping to target and off-target regions in tumor whole exome sequence data was corrected for GC content, target size, and sequence complexity and compared with a reference panel of normal samples to establish log copy ratio values as a measure of relative copy number across the genome (18). Bin-level copy ratio values were segmented using circular binary segmentation (19). Segment copy ratio values and minor allele frequency of germline heterozygous SNPs overlapping the segments were analyzed to determine the purity and ploidy of the sample, and allele-specific copy number for segments using an in-house pipeline. Next, we used SCHISM (20) to determine the cellular prevalence of mutations based on the observed variant allele frequency, estimated copy number, and sample purity by following an approach similar to that previously described (13). This approach to clonality assessment was not feasible for mutations in four cases (CGLU168, CGLU206, CGLU219, and CGLU249) where purity and ploidy could not be determined due to low tumor content. Mutation cellularity analysis is summarized in Supplementary Table S7.

TCR sequencing and differential expansion analyses

TCR clones were evaluated in pretreatment tumor tissue (with the exception of CGLU117, where tumor tissue from the time of resistance was also analyzed), and 40 serial peripheral blood lymphocytes (PBL) by next-generation sequencing in the metastatic NSCLC cohort (Supplementary Table S8). DNA from pretreatment tumor samples and PBLs was isolated by using the Qiagen DNA FFPE and Qiagen DNA Blood Mini Kit, respectively (Qiagen). TCR-β CDR3 regions were amplified using the survey (or deep) (PBLs) ImmunoSeq assay in a multiplex PCR method using 45 forward primers specific to TCR β gene segments and 13 reverse primers specific to TCR β gene segments (Adaptive Biotechnologies; refs. 21, 22). Productive TCR sequences were further analyzed. TCR sequencing data from TILs were used to identify tumor-specific TCR clonotypes in the peripheral blood. Peripheral TCR clones achieving a frequency of at least 0.005% were evaluated for differential abundance between baseline and the time of radiographic response using Fisher exact test with FDR P-value correction (corrected P ≤ 0.05). Those differentially abundant clones also found in the tumor were further selected to determine their frequencies in peripheral blood prior to treatment, at the time of response and upon emergence of resistance (Supplementary Tables S9 and S18). We calculated the average productive frequency of differentially abundant clones and used it as a metric of TCR dynamics during therapy. To cluster significantly expanded intratumoral TCR-β CDR3s based on potential recognition specificity, we used the GLIPH method (Grouping of Lymphocyte Interactions by Paratope Hotspots; ref. 23).

CDR3 and V1 gene usage analyses

Subsequent to initial filtering, we further reduced noise by eliminating clones that did not have frequencies beyond a mean rate of five counts. Thus, when two points were examined (baseline and time of radiologic response), the total sum of counts were greater or equal to 10. Using these data, we examined the usage of CDR3b Variable (V) and joining (J) regions, and their overall clonal composition by known significant clones at the two time points.

Multiplex cytokine immunoassay

We used a multiplex bead-based immunoassay on the Luminex platform that examines cytokines involved in T-cell activation, expansion, differentiation, and long-term proliferation (IFNy, IL1, IL2, Th1 immune response (IL12), acquisition of the Th2 phenotype (IL4) as well as immunosuppressive cytokines important for regulatory T cells (IL10) in four patients of the early stage.
cohort where additional serum was available. Differences in concentration of cytokines were evaluated between baseline and on treatment (week 2–6) samples.

Statistical analyses

cDNA values were dichotomized as detectable and undetectable. Characteristics for each group were compared using chi-square or Fischer exact test for categorical variables. Pearson correlation coefficient (R) was used to assess correlations between continuous variables. Differences between molecular responders and nonresponders were assessed by the Mann–Whitney test. Tumors were classified based on their nonsynonymous sequence alteration load in high and low mutators as previously described (24). The median point estimate and 95% confidence interval for PFS and OS were estimated by the Kaplan–Meier method. Survival curves were compared by using the log-rank test. Univariate Cox proportional hazards regression analysis was used to determine the impact of cDNA molecular response on PFS and OS. All P values were based on two-sided testing and differences were considered significant at P < 0.05. Statistical analyses were done using the SPSS software program (version 25.0.0 for Windows; IBM).

Results

Overall approach and patient characteristics

We analyzed 105 serial blood samples from 38 patients with NSCLC, including 24 patients with metastatic NSCLC during immune checkpoint blockade and 14 patients with stage I to IIIA surgically resectable NSCLC that received anti-PD1 therapy as part of a clinical trial of neoadjuvant nivolumab (Table 1; Supplementary Table S1; ref. 15). The median duration of follow-up was 12.7 months (range 3.0–37.8 months) and 16 months (2 years) for the metastatic and early-stage patients, respectively, and median duration of treatment was 7 months (range 1–20 months) for the metastatic cohort. We evaluated response to immune checkpoint blockade using standard computed tomographic (CT) imaging and changes in tumor burden were assessed by RECIST 1.1. Blood samples for the metastatic patients with NSCLC were prospectively collected prior to therapy, at an early time point between 4 and 8 weeks from treatment initiation and at additional serial time points during therapy until the time of disease progression (Supplementary Tables S2 and S3). For the patients with early-stage NSCLC treated with anti-PD1 therapy in the neoadjuvant setting, blood samples were collected prior to immunotherapy, at 2 weeks, immediately prior to resection and post-resection (Supplementary Table S2). cDNA was measured using the TEC-Seq approach (14) and the TCR repertoire was studied longitudinally by means of TCR sequencing (Fig. 1). Given the possibility of hematopoietic alterations that may be detected in the plasma (14), especially in heavily treated patients, we focused only on tumor-specific alterations. cDNA dynamics reflected clonal evolution under selective pressure of anti-PD1 therapy and emergence of immune escape. In such cases, tumor-specific variants were undetectable at the time of response followed by increase in mutant allele fraction at the time of acquired resistance (Supplementary Fig. S1). Emergence of molecular resistance preceded disease progression on imaging by an average of 10.8 weeks. Overall, cDNA-based molecular responses were detected on average 8.7 weeks earlier than conventional RECIST 1.1 response assessment (6.7 weeks vs. 15.4 weeks, P = 0.004; Supplementary Fig. S3).

Early cDNA clearance was a significant prognostic factor for PFS and OS. Patients with a reduction of cDNA to undetectable levels demonstrated a significantly longer PFS and OS compared with patients with no evidence of cDNA elimination (log-rank test, P = 0.001 and 0.008, respectively; Fig. 4A and B; Supplementary Fig. S4). The duration of the molecular responses tightly correlated with PFS and OS (Supplementary Fig. S5).

Radiographic imaging at the time of first assessment was a worse predictor of outcome to anti-PD1 therapy compared with cDNA molecular response for these patients (Fig. 4A and B; Supplementary Fig. S6). Patients with radiographically stable disease (n = 12) had differential responses to immune checkpoint blockade that were consistent with their molecular response pattern (Supplementary Fig. S7). More specifically, five patients with stable disease by imaging showed a clear molecular response pattern, with cDNA elimination between week 4 and 13 from immune checkpoint blockade initiation (Supplementary Fig. S1). All five patients derived clinical benefit from PD-1 blockade (PFS and OS ranging from 7.3 to 13.6 and 12 to 21.3 months, respectively; Supplementary Fig. S7), suggesting that imaging failed to detect the magnitude of therapeutic response. Interestingly, cDNA molecular responses more accurately predicted PFS and OS compared with tumor mutation burden in our
Table 1. Summary of clinical and molecular characteristics

<table>
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<tr>
<th>Patient</th>
<th>Immune-targeted therapy</th>
<th>Metastatic vs. early stage</th>
<th>Progression-free survival (0: progression; 1: progression) PFS (months)</th>
<th>Overall survival (0: censored; 1: dead of disease) OS (months)</th>
<th>Number of tumor-specific variants at baseline</th>
<th>ctDNA molecular response (0 = no, 1 = yes)</th>
<th>Clonal TCR expansion (0 = no, 1 = yes)</th>
<th>Tumor mutation burden</th>
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**NOTE:** Major pathologic response (MPR) was defined as <10% viable tumor cells at the time of surgical resection (15). TCR expansion was assessed at the time of radiographic response.

Abbreviations: PFS, progression-free survival; OS, overall survival; ND, not detected; N/A, not evaluable.

*Exome sequencing was not performed; for these cases, CLIA-targeted NGS was performed for clinical purposes.

*The baseline tumor was not available for whole exome sequencing; the resection sample was analyzed and used to identify tumor-specific variants in ctDNA.
cohort (TMB; Supplementary Fig. S8). When TMB and ctDNA were combined the ctDNA-based molecular responders clustered together independent of the TMB for both PFS and OS (Fig. 4C and D). Given that clonal mutation burden may be a more accurate predictor of response to immune checkpoint blockade, we performed survival analyses incorporating clonal TMB and we again found that ctDNA dynamics predict survival independent of clonal TMB status (Supplementary Fig. S9).

Molecular responses predict pathologic response to immune checkpoint blockade

Given the challenges with radiologic response assessments to immune checkpoint blockade, we sought to validate our observations in a NSCLC cohort where the therapeutic effect was measured at a pathologic level instead of using conventional imaging. We analyzed serial plasma samples from a recently reported clinical trial of neoadjuvant nivolumab for early-stage operable NSCLC (15). For these patients, the therapeutic effect was rigorously measured by pathologic assessment of residual tumor after two doses of anti-PD1 therapy (15, 25). Similar to our initial analyses, we observed that all tumors with a major or partial pathologic response to anti-PD1 therapy demonstrated a molecular response pattern of elimination of tumor-specific mutations in the circulation (Fig. 5A; Supplementary Fig. S10). In contrast, tumors without a pathologic response demonstrated a molecular resistance...
Peripheral TCR landscape and therapeutic outcome

We investigated how immune checkpoint blockade affects the peripheral TCR repertoire and whether there are TCR clonotype dynamic changes reflective of a systemic antitumor immune response. We focused our analyses on TCR clones found in the tumor microenvironment using TCR sequencing and investigated their dynamics in the peripheral blood, identifying those with a statistically significant differential abundance from baseline. Twelve of the 24 metastatic patients with NSCLC had available samples from both tumor infiltrating lymphocytes and peripheral blood lymphocytes for analysis (Supplementary Tables S3 and S8), including five that had previously undergone TCR sequencing (13) but had not been analyzed using this approach.

Similar to ctDNA analyses, we observed distinct patterns in TCR clonotype dynamics among the analyzed patients. For patients with clinical responses to immune checkpoint blockade, a statistically significant oligoclonal expansion of preexisting intratumoral T-cell clones was observed in peripheral blood at the time of radiologic response to PD1 blockade (CGLU111, CGLU117, CGLU127, and CGLU212; Fig. 2; Supplementary Fig. S11; Supplementary Tables S9–S12). For patients who developed acquired resistance, productive frequencies of intratumoral clones significantly decreased in peripheral blood at the time of acquired resistance (CGLU117, CGLU127, CGLU135, and CGLU161; Supplementary Fig. S11; Supplementary Tables S10, S11, S13, S14), with a timing that was similar to ctDNA analyses for most cases.

In contrast, for patients CGLU121 and CGLU115 that had primary resistance to immunotherapy, we did not identify any differentially abundant TCR clones among serial peripheral blood samples (Fig. 3; Supplementary Fig. S12). These patients progressed radiographically within 5 to 13 weeks from initiation of therapy and, in line with the clinical course, there was no evidence of TCR clonal expansion among the intratumoral TCR repertoire. A transient oligoclonal TCR expansion was observed for nonresponding patient CGLU159 at week 11; however, productive frequencies of differentially abundant clones quickly decreased to baseline levels at week 16, which coincided with disease progression (Supplementary Fig. S12; Supplementary Table S15). Patients CGLU162, CGLU1203, and CGLU243 had 1 to 26 intratumoral TCR clones with differential abundance at the time of best radiographic response compared with baseline but were classified as ctDNA molecular nonresponders (Supplementary Fig. S12; Supplementary

Figure 2. ctDNA and TCR clonal dynamics for a patient with sustained response to anti-PD1. A, ctDNA (TP53 993+1G>T mutation, blue) decreased to undetectable levels, signifying a complete molecular response at week 4. In contrast, CT imaging did not accurately capture the rate (B) or timing (C) of tumor regression (RECIST tumor burden dynamics, green). A complete response by RECIST 1.1 was achieved 26 weeks later than the molecular response (C). In parallel, TCR repertoire dynamics revealed clonotypic amplifications of intratumoral TCR clones in peripheral blood at the time of radiographic response. TCR clones with statistically significant differential abundance were evaluated as individual clones (D) and as a composite of productive frequencies (E). Patient was off anti-PD1 therapy and on immunosuppressive therapy at week 30 (arrow) due to emergence of immune-related toxicity.
Tables S16–S18. Patients CGLU203 and CGLU243 had unfavorable outcome to anti-PD1 therapy, suggesting that for these patients, ctDNA kinetics may more accurately predict therapeutic outcome.

We did not identify any shared TCR clones among the differentially expanded ones for all patients analyzed, consistent with the notion that the mutation-associated neoantigen repertoires are largely private. We evaluated putative shared CDR3 motifs among significantly expanded TCR clones using the grouping of lymphocyte interactions by paratope hotspots algorithm (23). Interestingly, TCR clones CSARVGVGNTIYF and CSARSGVGNTIYF, which were differentially abundant at the time of response to immune checkpoint blockade for patient CGLU127 and CGLU135, respectively, clustered together, suggesting a common specificity to a tumor- or mutation-associated antigen. We subsequently investigated potential differential sequence features focusing on Variable (V) and Joining (J) gene usage and CDR3 lengths among different timepoints for each patient. Usage of specific V and J gene segments increased at the time of response compared with baseline for a patient with sustained response (CGLU111) in contrast to a representative patient with primary resistance (CGLU112; Supplementary Fig. S13). Our findings on differential V gene usage may suggest clonotypic amplifications of specific immune subsets (CD8+ vs. CD4+) during immune checkpoint blockade (26).

**Discussion**

The unique nature of responses to immune checkpoint blockade (27, 28) and known limitations of conventional radiologic response assessments (3) highlight the need for development of biomarker-driven approaches to interpret therapeutic responses. Success of immunotherapy approaches depends on choosing patient populations most likely to benefit. There is therefore an urgent clinical need for molecular assays of response and resistance to immune-targeted agents. To this end, we analyzed ctDNA and TCR clonal dynamics during immune checkpoint blockade in NSCLC and assessed the value of longitudinal monitoring of liquid biopsies as a surrogate for response to therapy. Our findings indicate that ctDNA dynamics after treatment initiation may allow patients with primary resistance to immune checkpoint blockade to be rapidly identified and redirected to receive alternative options.

Noninvasive detection and monitoring of acquired resistance to EGFR-targeted therapy has been evaluated by serial sampling of ctDNA (29, 30) and as we have shown in a complementary study, changes in ctDNA levels may predict response to targeted therapy in NSCLC (8). Longitudinal assessment of ctDNA in patients with metastatic melanoma receiving anti-PD1 therapy has been demonstrated to be an accurate predictor of tumor response and therapeutic outcome (31) and early ctDNA clearance may correlate with durable clinical benefit to PD-1 blockade (10, 12).
ctDNA dynamics may be also informative in differentiating pseudoprogression from disease progression during immunotherapy (32). However, these approaches have been limited by low sensitivity and specificity of ctDNA methods, with many patients lacking detectable alterations and potential admixture between tumor alterations and those involved in clonal hematopoiesis (33, 34).

Moreover, interpretation of ctDNA analyses without knowledge of tumor-specific somatic alterations may be difficult in the setting of heavily pretreated patient populations such as patients with late-stage lung cancer, given the mutagenic effects of systemic chemotherapy and ionizing radiation on cells of the myeloid lineage (35). To address the possible presence of alterations in cfDNA from clonal nonmalignant hematopoietic cells, we have focused our analyses of variants in ctDNA that were also identified through next-generation sequencing of the matched tumor, allowing distinction of tumor-specific from blood cell proliferation variants.

Clonal expansion of intratumoral T cells may predict therapeutic outcome for immune checkpoint blockade (36); however, little is known about the significance of peripheral expansion of TCR clones found in the tumor microenvironment during therapy. Expansion of peripheral CD8$^+$ T-cell populations has been shown to precede immune-related adverse events in patients treated with ipilimumab (37). We investigated whether the patients with NSCLC in our cohort developed immune-related adverse events at the time of TCR clonal expansion and did not identify a definitive pattern with the exception of patient CGLU243, where pneumonitis emerged shortly after treatment initiation. Although there were cases for which TCR expansion preceded the development of a grade 2 to 4 immune-related adverse event (CGLU161, CGLU117), such events were also noted significantly later from the time of TCR expansion (CGLU111, CGLU135). These observations highlight the challenges with interpretation of the evolving peripheral TCR repertoire. Assessing the quality of the immune

Figure 4.
Early ctDNA clearance predicts PFS and OS. Patients with reduction of ctDNA to undetectable levels demonstrated a significantly longer PFS and OS compared with patients with no evidence of ctDNA elimination (log-rank $P = 0.001$ and 0.008, respectively; A and B). Patients with undetectable ctDNA (molecular responders) clustered together independent of their tumor mutation burden (C) and the same pattern was observed for patients with detectable ctDNA (molecular nonresponders; D). Patients with ctDNA molecular response and either high or low tumor mutation burden had a significantly longer PFS and OS (log-rank $P = 0.015$ and 0.027, respectively).
whether a liquid biopsy approach can replace conventional radiologic progression will improve outcome and ultimately whether switching therapy based on ctDNA dynamics prior to tumor mutations. Prospective studies will be needed to address the frequency of serial monitoring and feasibility of subsequent therapies. Additional work will be needed to address the frequency of serial monitoring and feasibility of interpreting ctDNA dynamics without prior knowledge of tumor mutations. Prospective studies will be needed to address whether switching therapy based on ctDNA dynamics prior to radiologic progression will improve outcome and ultimately whether a liquid biopsy approach can replace conventional imaging as a gold standard for early response assessment to immune checkpoint blockade.

**Disclosure of Potential Conflicts of Interest**

P.M. Forde reports receiving a commercial research grant from AstraZeneca, BMS, Corvus, Kyowa, Merck, and Novartis, and is a consultant/advisory board member of Abbvie, AstraZeneca, BMS, Boehringer, EMD Serono, Lilly, Merck, and Novartis. J.R. White is a consultant at Personal Genome Diagnostics and has ownership interest (including stock, patents, etc.) in Respexa Biosciences. J. Naidoo reports receiving a commercial research grant from AstraZeneca/MedImmune and Merck, is a consultant/advisory board member of Bristol-Myers Squibb and is a consultant/advisory board member of Bristol-Myers Squibb and AstraZeneca/MedImmune. V. Adleff is a consultant/advisory board member of Personal Genome Diagnostics. C.L. Hann reports receiving other commercial research support from Bristol-Myers Squibb and is a consultant/advisory board member of AstraZeneca/MedImmune, Bristol-Myers Squibb, and Genentech. B. Levy is a consultant/advisory board member of Genentech, AstraZeneca, Eli Lilly, Celgene, and Takeda. J. Feliciano is a consultant/advisory board member of AstraZeneca, Merck, Genentech, and Eli Lilly, and has provided expert testimony for legal review. P. Illei reports receiving other commercial research support from Bristol-Myers Squibb and is a consultant/advisory board member of Bristol-Myers Squibb and Genentech. J. Taube reports receiving other commercial research support from Bristol-Myers Squibb and is a consultant/advisory board member of Bristol-Myers Squibb, AstraZeneca, Merck, and Amgen. D.M. Pardoll reports receiving a commercial research grant from Bristol-Myers Squibb and is a consultant/advisory board member of Bristol-Myers Squibb, AstraZeneca, Merck, and Amgen. J. Sauter has ownership interest (including stock, patents, etc.) in Lumacys, Inc. J.L. Sauter has ownership interest (including stock, patents, etc.) in Merck & Company, Inc. (New), Chemed Corporation, Pfizer, Thermo Fisher Scientific, Celgene, and Allergan PLC SHS. J. Tauke reports receiving a commercial research grant from Bristol-Myers Squibb and is a consultant/advisory board member of AstraZeneca, Compugen, Merck, has ownership interest (including stock, patents, etc.) in WindMil Therapeutics, Potenza, Erveax, Dracen Pharmaceuticals, DNAtrix, Aduro Biotech, Five Prime, and Tizona, and is a consultant/advisory board member of Amgen, Bayer, Camden Partners, Janssen, FLX Bio, Immunomics, and Rock Springs Capitol. J.E. Chat is a consultant/advisory board member of AstraZeneca, Merck, BMS, and Genentech. M.D.
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References

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Dynamics of Tumor and Immune Responses during Immune Checkpoint Blockade in Non–Small Cell Lung Cancer

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