Applications of circulating tumor DNA in a cohort of phase I solid tumor patients treated with immunotherapy

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Abstract

**Background:** The correlation between blood-based tumor mutation burden (bTMB) and tissue-based TMB (tTMB) has not been broadly tested in a multi-cancer cohort. Here, we assess the correlation between bTMB with tTMB in phase I trial patients treated with immunotherapy. As an exploratory analysis, we evaluated circulating tumor DNA (ctDNA) dynamics in responders.

**Methods:** Patients treated with immunotherapy at the Princess Margaret phase I trials unit were enrolled. Pre-treatment plasma ctDNA and matched normal blood controls were collected. Available archival tissue formalin-fixed paraffin-embedded (FFPE) samples were analyzed. A 425-gene panel was used to sequence both ctDNA and FFPE samples. Samples with TMB within the highest tertile were considered as high TMB.

**Results:** Thirty-eight patients were accrued from 25 different trials, 86.8% of which involved a PD-1/PD-L1 inhibitor. Thirty patients (78.9%) had detectable mutations in ctDNA, of which the median (range) bTMB was 5 (1-53) mut/Mb. Of the 22 patients with available FFPE samples, mutations were detected in 21 (95.4%); the median (range) tTMB was 6 (2-124) mut/Mb. Among the 16 patients with detectable mutations in both FFPE and ctDNA, a statistically significant correlation between bTMB and tTMB was observed ($\rho=0.71$; $p=0.002$). High TMB was not associated with better survival. All 3 responders had a decrease in the variant allele frequency (VAF) of mutations detected in ctDNA at a second time-point relative to baseline, indicating a potential early marker of response.

**Conclusions:** In this small series, bTMB correlated with tTMB. An on-treatment decrease in VAF of mutations detected in ctDNA at baseline was observed in responders. Larger studies to verify our findings are warranted.
Tissue-based tumor mutational burden (tTMB), or the total number of mutations per megabase of coding sequence in a tumor specimen, is a potential predictive marker for response to immunotherapy [1]. Increased tTMB has been associated with a higher likelihood of IMMUNOTHERAPY response[2]. Determination of tTMB was initially pioneered through whole exome sequencing (WES). However, several recent publications demonstrated that WES tTMB results correlate well with tTMB estimated by large (i.e. > 300 genes) Next Generation Sequencing (NGS) panels that are more routinely used in clinical practice[3, 4]. Nonetheless, many other challenges limit the application of tTMB as a selection biomarker for IMMUNOTHERAPY treatment, including logistical aspects such as tissue availability and turn-around time for results; sampling issues due to intratumoral heterogeneity; and technical aspects such as lack of standardization in NGS platforms, cut-offs and reproducibility[5].

Blood-based TMB (bTMB), the calculation of TMB through analysis of mutations detected in circulating tumor DNA (ctDNA), is an emerging alternative that may overcome some of the barriers associated with tTMB. Pre-treatment bTMB correlates well with tTMB in metastatic non-small cell lung cancer (NSCLC)[6, 7] and castrate-resistant prostate cancer[8]; however, this has not been broadly tested in other cancer types. Beyond bTMB calculation, targeted ctDNA panels may provide prognostic information as well as inform treatment decisions[9]. For instance, the identification of driver mutations (e.g. \textit{EGFR T790M} in NSCLC) can guide genotype-directed targeted therapy, whereas the identification of resistance mutations (e.g. \textit{EGFR C797S} mutation in NSCLC) can prevent futile treatments.

In this pilot study, we analyzed a cohort of advanced solid tumor patients undergoing immunotherapy treatment in an academic phase I clinical trials unit. We hypothesize that bTMB
and tTMB would be highly correlated, irrespective of tumor histology, in this heterogeneous pan-patient population. In addition, we postulated that phase I trial patients whose tumors harbor high bTMB would have a favorable clinical outcome in response to immunotherapy treatment. As an exploratory analysis, we evaluated ctDNA dynamics in patients who responded to immunotherapy treatment.

Material and Methods

Patients and samples

From December 2017 to July 2018, patients with metastatic solid tumors seen at the Princess Margaret Cancer Centre phase I trials unit and enrolled in an early phase clinical trial involving investigational immunotherapy treatment were included in this analysis. Investigational immunotherapy treatments such as immune checkpoint inhibitors, vaccines, cytokines, and oncolytic viruses (either as monotherapy, or in combination with other immunotherapy agents, or with molecular targeted agents) were included. Combinations involving chemotherapy were not included. Pre-treatment and on-treatment whole blood samples were collected in Streck tubes (Cell-Free DNA BCT®) and separated into plasma and buffy coat cell fractions in accordance with local standard operating procedures, via an institutional liquid biopsy program (“LIBERATE”, Liquid Biopsy Evaluation and Repository Development at Princess Margaret, NCT03702309). Circulating tumor (ct)DNA was extracted from plasma samples collected at pre-treatment, mid-cycle 1 (if feasible), pre-cycle 2, 3, and every other cycle thereafter, and at the end of treatment. Somatic alterations were filtered with matched germline DNA obtained from buffy coat
peripheral blood mononuclear cells to remove germline mutations. When available, matched archival formalin-fixed, paraffin-embedded (FFPE) tumor tissues were obtained for genomic characterization. Pre-treatment plasma ctDNA, germline DNA, and archival FFPE tumor tissues were assayed. In addition, for selected cases, additional on-treatment samples were assayed to evaluate ctDNA dynamics of patients that responded and not responded to immunotherapy. This study has ethical approval (18-5815).

**Sample extraction, library preparation and sequencing**

Whole blood was collected in Cell-Free (cf)DNA BCT tubes (Streck, USA). Plasma and buffy coat were isolated from whole blood after centrifugation at 1900g for 10 minutes. The plasma layer was further centrifuged at 16,000g for 10 minutes prior to cfDNA extraction with QIAamp Circulating Nucleic Acid kit (Qiagen, USA). Genomic DNA from FFPE tumor tissue and buffy coat was extracted with the AllPrep DNA/RNA FFPE kit and AllPrep DNA/RNA/miRNA Universal kit (Qiagen, USA) respectively. Purified DNA was quantified by Qubit 3.0 using the dsDNA HS Assay kit (ThermoFisher Scientific, USA).

A customized panel targeting 425 cancer-relevant genes — GeneseeqPrime™ — was used for hybridization enrichment (see the Supplementary Appendix, available online). The capture reaction was performed with Dynabeads M-270 (Life Technologies, CA, USA) and xGen Lockdown hybridization and wash kit (Integrated DNA Technologies) according to manufacturers’ protocols. Captured libraries were on-beads PCR amplified with Illumina p5 (5’ AAT GAT ACG GCG ACC ACC GA 3’) and p7 primers (5’ CAA GCA GAA GAC GGC ATA CGA GAT 3’) in KAPA HiFi HotStart
ReadyMix (KAPA Biosystems), followed by purification using Agencourt AMPure XP beads. Libraries were quantified by qPCR using KAPA Library Quantification kit (KAPA Biosystems). Library fragment size was determined by Bioanalyzer 2100 (Agilent Technologies). The target-enriched library was then sequenced on HiSeq4000 NGS platforms (Illumina) according to the manufacturer’s instructions. The average coverage depth was 5569X (2626-8701X), 1410X (963-2582X), and 354X (285-453X) for plasma, tumor, and normal control samples, respectively. The average Q30 base percentage was 93% for plasma samples, 88% for tumor samples, and 92% normal control samples. Detailed quality control results are presented in Supplementary Table 1.

**Mutation calling and TMB definition**

Trimmomatic[10] was used for FASTQ file quality control. Leading/trailing low quality (quality reading below 20) or N bases were removed. Pair-end reads were then aligned to the human reference genome (hg19) using Burrows-Wheeler Aligner (BWA)[11] with default parameters. PCR deduplication was performed using Picard V2.9.4 (Broad Institute, MA, USA). Local realignment around indels and base quality score recalibration were performed with the Genome Analysis Toolkit (GATK 3.4.0). Somatic single-nucleotide variants (SNVs) were identified using MuTect[12], and small insertions and deletions (indels) were detected using SCALPEL[13]. The cut-off for mutation detection was 1% somatic variant allele frequency (VAF) and 5 reads in plasma samples; 2% VAF and 5 reads in FFPE samples. For patients with multiple plasma samples, if a mutation meets the above cutoff in at least one sample, the detection cutoff for the same mutation was dropped in other samples to reduce false negatives. Final list of mutations was annotated using vcf2maf (available on github). TMB was counted by summing all base
substitutions and indels in the coding region of targeted genes, including synonymous alterations to reduce sampling noise and excluding known driver mutations as they are over-represented in the Panel, as previously described (24). Samples within the highest mutation-load tertile (top 33.3%) were classified as having high TMB (both for tTMB and bTMB). Discordance in bTMB and tTMB is defined when these values fall into different categories as defined by their respective cut-offs for high versus low TMB (e.g. high bTMB and low tTMB, low bTMB and high tTMB).

**Statistical analysis**

Correlations between tTMB and bTMB were calculated using Spearman’s rank test. Demographics characteristics were summarized in means, medians and proportions. Associations between categorical variables were examined using the Chi-square or Fisher’s exact test. For survival analyses, Kaplan-Meier curves were compared using the log-rank test, and hazard ratios (HRs) were calculated by Cox proportional hazards model. No formal power calculations were performed for this exploratory analysis. A two-sided p value of less than 0.05 was considered statistically significant. All statistical analyses were performed in R (v.3.3.2).

**Results**

**Patient characteristics**

Among the 40 patients enrolled, 2 were screen failures and did not receive immunotherapy treatment leaving 38 patients who received at least 1 dose of investigational immunotherapy. This cohort included 25 different phase I studies, and 28 tumor types mapped by the OncoTree
ontology[14]. After grouping, the most frequent tumor sites were colorectal, head and neck squamous cell carcinoma (HNSCC) and breast cancer, all with 5 patients (13.1%) each. The mean age was 59 years-old (range = 21 – 77) and 20 patients (52.6%) were female. Thirty-three patients (86.8%) received an anti-PD1/anti-PD-L1 immunotherapy treatment, and 31 patients (81.6%) participated in combination trials. Table 1 summarizes the demographic characteristics of the cohort. Baseline plasma sample from all 38 patients and archival FFPE samples from 22 patients (57.8%) underwent NGS targeted-sequencing (see the Consort diagram in Supplementary Figure 1).

Overall, 3 patients (7.8%) achieved partial response (PR), 11 patients (28.9%) achieved stable disease, and 24 patients (63.1%) had progressive disease as best response by RECIST v1.1. After a median follow-up of 15 months, the median progression-free survival (mPFS) for the entire cohort was 1.8 months (95% CI = 1.76- 1.92), and the median overall-survival (mOS) was 9.89 months (95% CI = 7.40 – 12.40); the 1-year PFS rate was 13.2% (95% CI = 5.8% - 29.8%) and 1-year OS rate was 39.5% (95% CI = 26.6% -58.5%).

**Correlation between bTMB and tTMB and association with immunotherapy efficacy**

We detected somatic mutations in 30 of 38 (78.9%) baseline plasma samples and 21 of 22 (95.4%) FFPE samples. The median bTMB was 5 (range = 1-53) mutations per megabase (mut/Mb) and tTMB was 6 (range = 2-124) mut/Mb. Sixteen patients had mutations detected in both plasma and FFPE samples. The Spearman TMB correlation of these 16 pairs was 0.71 (p = 0.002, Figure 1A). When stratified by top tertile TMB (12 mut/Mb for both sample types, when only these 16
pairs are assessed), 14 of 16 (87.5%) patients have concordant TMB status in both samples. The 2 patients with discordant TMB status had esophageal adenocarcinoma (pt0002) and microsatellite instability-high (MSI-H) colorectal cancer (pt0038), and both had higher tTMB than bTMB: 14 vs. 1 mut/Mb and 124 vs. 6 mut/Mb respectively, as well as stable disease (SD) as best RECIST 1.1 response. Furthermore, both patients were treated with chemotherapy in the interval between FFPE and blood collection acquisition, and pt0038 had two distinct POLE variants in the FFPE tissue sample, which were not detected in ctDNA. The median time interval between blood and FFPE tissue collection dates was 20 months (range = 6.9 – 64.3 months). The time-interval between blood and FFPE tissue collection was not correlated with the difference in TMB between sample types (Spearman correlation $\rho = 0.19$, $p = 0.48$, Figure 1B). The time interval difference for the two patients with discordant TMB results was 60.8 months for pt0002, and 15.6 months for pt0038 respectively.

Next, we performed a sensitivity analysis of the bTMB and tTMB correlation including all patients with matched samples available (22 pairs), regardless of mutation detection. For those subjects with samples without detectable mutations, a TMB of 0 (zero) was considered. The Spearman correlation was $\rho = 0.48$ ($p = 0.02$, Figure 1C). When stratified by top-tertile TMB (5 mut/Mb for bTMB, and 12 mut/Mb for tTMB, when 22 pairs are assessed), there were 3 patients with discordant tTMB and bTMB: pt0002 (described previously), pt0016 (a 61-year-old male with metastatic HNSCC with a tTMB of 55 and a bTMB of 0, who had PD as best response) and pt0045 (a 49-year-old woman with metastatic triple negative breast carcinoma with a tTMB of 9 and a bTMB of 7, who had PR as best response). The time-interval between blood and FFPE tissue collection dates for patients pt0016 and pt0045 was 13.8 and 21.9 months, respectively.
Subsequently, we examined whether bTMB or tTMB status was associated with survival or overall response rate. Using a cutoff of 12 mut/Mb, both PFS and OS did not statistically significantly differ between low TMB and high TMB group, regardless of sample types (Supplementary Figure 2A-D). However, the 3 PR patients had higher median bTMB compared to both patients with SD (45 vs 1) and PD (45 vs 5, Figure 1D).

cTDT DNA mutational landscape and sequential cTDT DNA reduction in PR patients

Overall, somatic mutations were identified in 156 genes, and 14 genes were encountered in more than 3 patients (Figure 2). The most frequently mutated genes were TP53 in 16 patients (53.3%), NOTCH2 in 5 patients (16.6%), and PKHD1 in 5 patients (16.6%).

To monitor the dynamic changes in the variant allele frequency (VAF) of specific mutations detected in plasma samples of the 3 PR patients, we sequenced additional samples collected at the following time points: cycle 2 (typically one month from baseline), best response, and latest available timepoint. Sequencing results revealed that VAF of mutations detected in the baseline samples were reduced at the second timepoint and stayed minimally detectable through the treatment course (Figure 3A-C). More specifically, pt0013, a 70 year-old man with metastatic anal squamous cell carcinoma with disease involving lungs, liver, lymph nodes and peritoneum treated with a combination involving an anti-PD1/PDL1 antibody as first line regimen had 48 non-synonymous mutations present in his baseline plasma sample, 16 of which had VAF over 10%. At the second time point, only 9 mutations were detectable and were all less than 1% VAF. Pt0022, a 50 year-old woman with metastatic MSI-H endometrial carcinoma with loco-regional
recurrence and lymph node metastasis, treated with a combination involving an anti-PD1/PDL1 agent as second-line regimen had 36 non-synonymous mutations present in her baseline plasma sample, with VAF between 1% and 9%. The number of detectable mutations decreased to 25 at the second timepoint, all with less than 1% VAF. Lastly, pt0045, a 49-year-old woman with metastatic triple negative breast cancer involving lungs and lymph nodes on second-line treatment with a combination involving an anti-PD1/PD-L1 agent had 7 somatic mutations present in her baseline plasma sample, with VAF between 2% and 9%. In the second sample all 7 mutations’ VAF were reduced to less than 1%.

Next, we sequenced samples of 5 randomly selected patients who had PD as best response (pt0005, a 60-year-old male with metastatic pancreatic neuroendocrine tumor; pt0012, a 76-year-old female with metastatic cutaneous melanoma; pt0015, a 44-year-old female with metastatic triple negative breast cancer; pt0051, a 65-year-old male with metastatic duodenal adenocarcinoma; and pt0052, a 51-year-old male with metastatic small cell lung cancer). Of these, only pt0005 had an on-treatment decrease of the VAF of ctDNA detectable mutations at baseline (Supplementary Figure 3A), whereas the remainder 4 patients had either on-treatment stability or increase of the VAF of ctDNA detectable mutations at baseline (Supplementary Figure 3B-E).

**Discussion**

Our results demonstrate a positive correlation between bTMB and tTMB in a multi-cancer cohort treated with immunotherapy as part of an early phase clinical trial program at the Princess...
Margaret Cancer Centre. The correlation coefficient of $\rho = 0.71$ ($p = 0.002$) found in patients with both blood and tissue samples containing detectable mutations in our study is in agreement with the correlation between tTMB and bTMB identified in NSCLC[6, 7]. Moreover, when using top tertile as a benchmark for defining high TMB (12 mut/Mb for both bTMB and tTMB), only 2 patients had discordant results. Differences between time of blood collection and FFPE acquisition could not explain discrepancies between tTMB and bTMB results in our cohort. However, both patients with discordant results received chemotherapy treatment in the interval between tissue and blood collections, which can potentially account for some of the reduction in bTMB compared to tTMB. A sensitivity analysis including all patients with available samples regardless of mutation detection (22 pairs) demonstrated a positive, but weaker correlation coefficient of $\rho = 0.48$ ($p = 0.02$) suggesting that bTMB and tTMB correlation is better when mutations are detected in both sample types. Multiple factors may limit tTMB and bTMB correlation in our study, including heterogeneity of the cohort comprising multiple tumor types with different genomic alterations; differences in the sensitivity of variant detection between FFPE and plasma; variations in tumor cellularity; temporal and spatial heterogeneity of samples; treatment effects, etc. Nevertheless, our results suggest that bTMB may be a potential substitute for tTMB for the majority of patients whose tumors harbor detectable mutations in ctDNA.

High TMB (top tertile) assessed by tTMB or bTMB was not associated with better survival outcomes in the current cohort. These findings are similar to data presented by Marabelle et al. of a 755 patients multi-cancer cohort treated with single agent pembrolizumab[15]. High tTMB, defined as $\geq 10$ mut/Mb in that study, was not associated with improved survival, although patients with high tTMB were more likely to respond to immunotherapy treatment[15].
Nonetheless, these findings, akin to our results, may reflect the heterogeneity of pan-cancer patient cohorts and are underpowered to assess the prognostic value of TMB in a tumor-specific context. TMB varies accordingly to cancer type[2], thus different tumor types likely require distinct cut-off values. A universal cut-off value may not be appropriate in pan-cancer analyses. Samstein et al., analyzing genomic data of 1662 patients who underwent NGS targeted sequencing (MSK-IMPACT, with 468 genes) and received an immunotherapy-based treatment, found that within the same histology, the higher the tTMB, the higher the likelihood of better survival outcomes[16]. In this MSK-IMPACT study, the highest mutation-load quintile (top 20%) for each tumor type was selected as the cut-off to define high TMB; and the authors found that different tumors yield different cut-offs, e.g. breast 5.9 mut/Mb and melanoma 30.7 mut/Mb. Due to our limited sample size and few patients with objective responses (3 PRs), we have not performed analysis per histology. However, pt0045 in our cohort, a patient with metastatic triple negative breast cancer whose blood sample had a bTMB of 7 mut/Mb — typically classified as low, but considered high in the context of breast cancer — responded to immunotherapy treatment, supporting the histology-based TMB cut-off hypothesis. Nevertheless, further research is needed to translate the application of TMB into clinical practice and confirm its clinical utility, as this biomarker has been the subject of pronounced scrutiny[3, 5]. Some of the challenges lie in the lack of TMB definitions, standardization of assays, variant filtering methods, sequencing technologies, cut-offs and reporting of TMB results[17]. Working groups have been established in an attempt to harmonize some of these aspects[18].

Mutations in ctDNA were observed in 79% of patients assayed with the GeneseeqPrime™ panel in this cohort, which is comparable to the sensitivity of other commercially available ctDNA
panels testing a similar number of genes [6, 19, 20]. The ctDNA VAFs of all 3 responders had decreased in the blood sample collected at cycle 2 relative to baseline sample, whereas 4 of 5 patients with PD had either on-treatment stability or increase in ctDNA VAF of detectable mutations. One patient with discordant bTMB versus tTMB results with PD as best response had on-treatment decrease in ctDNA VAF of detectable mutations. In contrast to the 3 patients with PR as best response whose on-treatment ctDNA mutations had VAF of less than 1%, most of the PD patient’s on-treatment ctDNA mutations had VAF of greater than 1%. There is a general association between cfDNA mutation VAF clearance and treatment responses in different tumor types[21, 22]. For instance, in a cohort of patients with metastatic PIK3CA mutated breast cancer treated with fulvestrant and palbociclib, a post treatment decrease in plasma ctDNA PIK3CA levels below the median at cycle 1 day 15 compared to baseline levels, was associated with improved PFS[22]. Nevertheless, many factors determine the ability of a targeted panel to detect ctDNA mutations over time. Foremost, there is usually a trade-off between panel size and reading depth. For instance, applicability of WES in ctDNA is currently limited due to shallow coverage, and thus may not be able to track mutations that exist at very low levels. Conversely, despite having an adequate reading depth, small sized panels are applicable mainly in scenarios where the genomic regions of interest are known or anticipated (e.g. BRAF mutation in BRAF mutant melanomas), but are not comprehensive enough to be useful in an unselected scenario. A potential solution is the emergence of bespoke small-sized ctDNA panels with great read depths targeting mutations identified from WES (from pre-treatment tumor samples), which are highly sensitive for ctDNA detection. This type of bespoke ctDNA analysis was recently applied by our group in a multi-cancer cohort treated with single agent pembrolizumab[23, 24], dynamic
changes in the VAFs of patient-specific ctDNA samples collected on-treatment compared to baseline predicted for treatment response, PFS and OS. Our current data suggest that a large ctDNA panel may also be able to provide early response markers of immunotherapy treatment effectiveness. While the coverage depth of large ctDNA panels is inferior to a bespoke approach, large panels have the advantages of being readily available, potentially more scalable and not reliant on access to tumor WES results. In addition, large ctDNA panels have the potential advantage of capturing molecular progression caused by emergence of new mutations not previously detected at baseline[25].

This study has several limitations and results should be interpreted with caution. First, our cohort is small and includes patients with multiple tumor types and distinct genomic alterations. Second, patients receiving treatment across 21 different trials were included; nevertheless, 87% of patients received an anti-PD1/PD-L1 checkpoint inhibitor as part of their treatment. Third, 42% of patients had no available archival tissue, preventing a more definitive analysis of tTMB/bTMB correlation. Lastly, our small sample size prohibited multivariable adjustments.

In conclusion, our work demonstrates that when mutations are detected in both tissue and blood samples, tTMB and bTMB are highly correlated in a diverse multi-tumor phase I cancer clinical trials patient cohort. Blood-based TMB may be an alternative to tTMB in patients with advanced solid tumors with detectable mutations in ctDNA. An exploratory analysis suggests that early decrease in VAF of ctDNA mutations may be a marker of immunotherapy response. Larger studies addressing these hypotheses are warranted.
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**Author Contributions:** DVA and LLS conceived the study. AW performed data analysis. DVA reviewed clinical data and performed data analysis. AS, PLB, ARH, AAR and LLS accrued patients and supervised biospecimen collection. TJP and DT supervised biospecimen sample processing.
by AL and KM. HC and EP provided logistical support to the project. AM and HB performed the pathology review of samples. AW, HB and XW conducted the sequencing assays and provided technical expertise and scientific feedback. AW and LW conducted the statistical analysis. DVA, AW, XW and LLS wrote the manuscript. All authors discussed the results and implications and commented on the manuscript at all stages.

Data Availability

The data underlying this article cannot be shared publicly due to confidentiality agreements with sponsors of ongoing/completed Phase I studies from Princess Margaret Cancer Centre. This is a deidentified secondary analysis. The data may be shared upon reasonable request to the corresponding author.

References

1. Havel JJ, Chowell D, Chan TA. The evolving landscape of biomarkers for checkpoint inhibitor immunotherapy. Nature Reviews Cancer 2019;19(3):133-150.

2. Yarchoan M, Hopkins A, Jaffee EM. Tumor Mutational Burden and Response Rate to PD-1 Inhibition. New England Journal of Medicine 2017;377(25):2500-2501.

3. Chan TA, Yarchoan M, Jaffee E, et al. Development of tumor mutation burden as an immunotherapy biomarker: utility for the oncology clinic. Annals of Oncology 2018;30(1):44-56.
4. Fang W, Ma Y, Yin JC, et al. Comprehensive Genomic Profiling Identifies Novel Genetic Predictors of Response to Anti–PD-(L)1 Therapies in Non–Small Cell Lung Cancer. Clinical Cancer Research 2019;25(16):5015.

5. Addeo A, Banna GL, Weiss GJ. Tumor Mutation Burden—From Hopes to Doubts. JAMA Oncology 2019;5(7):934-935.

6. Gandara DR, Paul SM, Kowanetz M, et al. Blood-based tumor mutational burden as a predictor of clinical benefit in non-small-cell lung cancer patients treated with atezolizumab. Nature Medicine 2018;24(9):1441-1448.

7. Wang Z, Duan J, Cai S, et al. Assessment of Blood Tumor Mutational Burden as a Potential Biomarker for Immunotherapy in Patients With Non–Small Cell Lung Cancer With Use of a Next-Generation Sequencing Cancer Gene Panel. JAMA Oncology 2019;5(5):696-702.

8. Qiu P, Poehlein CH, Marton MJ, et al. Measuring Tumor Mutational Burden (TMB) in Plasma from mCRPC Patients Using Two Commercial NGS Assays. Scientific Reports 2019;9(1):114.

9. Araujo DV, Bratman SV, Siu LL. Designing circulating tumor DNA-based interventional clinical trials in oncology. Genome Medicine 2019;11(1):22.

10. Bolger AM, Lohse M, Usadel B. Trimmomatic: a flexible trimmer for Illumina sequence data. Bioinformatics 2014;30(15):2114-20.

11. Li H, Durbin R. Fast and accurate short read alignment with Burrows-Wheeler transform. Bioinformatics 2009;25(14):1754-60.

12. Cibulskis K, Lawrence MS, Carter SL, et al. Sensitive detection of somatic point mutations in impure and heterogeneous cancer samples. Nat Biotechnol 2013;31(3):213-9.
13. Fang H, Bergmann EA, Arora K, et al. Indel variant analysis of short-read sequencing data with Scalpel. Nat Protoc 2016;11(12):2529-2548.

14. Oncotree - http://oncotree.mskcc.org/#/home.

15. Marabelle A, Fakih MG, Lopez J, et al. Association of tumour mutational burden with outcomes in patients with select advanced solid tumours treated with pembrolizumab in KEYNOTE-158. Annals of Oncology 2019;30(Supplement_5).

16. Samstein RM, Lee C-H, Shoushtari AN, et al. Tumor mutational load predicts survival after immunotherapy across multiple cancer types. Nature Genetics 2019;51(2):202-206.

17. Fumet J-D, Truntzer C, Yarchoan M, et al. Tumour mutational burden as a biomarker for immunotherapy: Current data and emerging concepts. European Journal of Cancer 2020;131:40-50.

18. Stenzinger A, Allen JD, Maas J, et al. Tumor mutational burden standardization initiatives: Recommendations for consistent tumor mutational burden assessment in clinical samples to guide immunotherapy treatment decisions. Genes, Chromosomes and Cancer 2019;58(8):578-588.

19. Müller JN, Falk M, Talwar J, et al. Concordance between Comprehensive Cancer Genome Profiling in Plasma and Tumor Specimens. Journal of Thoracic Oncology 2017;12(10):1503-1511.

20. Khagi Y, Goodman AM, Daniels GA, et al. Hypermuted Circulating Tumor DNA: Correlation with Response to Checkpoint Inhibitor–Based Immunotherapy. Clinical Cancer Research 2017;23(19):5729.
21. Syeda MM, Wiggins JM, Corless B, et al. Circulating tumor DNA (ctDNA) kinetics to predict survival in patients (pts) with unresectable or metastatic melanoma treated with dabrafenib (D) or D + trametinib (T). Journal of Clinical Oncology 2019;37(15_suppl):9510-9510.

22. O’Leary B, Hrebien S, Morden JP, et al. Early circulating tumor DNA dynamics and clonal selection with palbociclib and fulvestrant for breast cancer. Nature Communications 2018;9(1):896.

23. Iafolla MAJ, Yang C, Dashner S, et al. Bespoke circulating tumor DNA (ctDNA) analysis as a predictive biomarker in solid tumor patients (pts) treated with single-agent pembrolizumab (P). Journal of Clinical Oncology 2019;37(15_suppl):2542-2542.

24. Yang C, Iafolla MA, Dashner S, et al. Bespoke circulating tumor DNA (ctDNA) analysis as a predictive biomarker in solid tumor patients (pts) treated with single agent pembrolizumab (P). Annals of Oncology 2019;30(Supplement_5).

25. Jamal-Hanjani M, Quezada SA, Larkin J, et al. Translational Implications of Tumor Heterogeneity. Clinical Cancer Research 2015;21(6):1258.
## Tables

### Table 1 – Demographic characteristics patients enrolled (N=38)

| Demographics                        | No. (%) |
|-------------------------------------|---------|
| Mean age (range), y                 | 59 (21-77) |
| Female                              | 20 (52.6) |

| Tumor Site                          |         |
|-------------------------------------|---------|
| Colorectal                          | 5 (13.1) |
| HNSCC                               | 5 (13.1) |
| Breast                              | 5 (13.1) |
| Other*                              | 23 (60.5) |

| Treatment                           |         |
|-------------------------------------|---------|
| Involves Anti-PD-1/PD-L1 Antibodies| 33 (86.8) |
| Combination Trial                   | 31 (81.6) |
| Prior Anti-PD1/PDL1 Antibodies      | 9 (23.7) |

| Response Rate by RECIST v1.1        |         |
|-------------------------------------|---------|
| Partial Response                    | 3 (7.8) |
| Stable Disease                      | 11 (28.9) |
| Progressive Disease                 | 24 (63.1) |

*Others (grouped): Esophagus; Ovary; NET; Cholangiocarcinoma; Renal cell carcinoma; Melanoma; ASCC; Endometrial; Mesothelioma; Prostate; Pancreas; Sarcoma; Germ cell tumor; Small Bowel; Small cell lung cancer. HNSCC = Head and Neck Squamous Cell Carcinoma.
**Figure Legends**

**Figure 1.** Correlations between bTMB and tTMB; the time interval between samples acquisition and differences in TMB; and the relationship between bTMB and overall response rate.

A) Spearman correlation of bTMB and tTMB in patients with detectable mutations in both samples (16 pairs). Dashed lines = top tertile TMB (12 mut/ Mb for both). B) Difference between bTMB and tTMB vs time interval between plasma and tissue acquisition. C) Spearman correlation of bTMB and tTMB in all patients with matched samples, regardless of mutation detection (22 pairs). Dashed lines = top tertile TMB (5 mut/Mb for bTMB and 12 mut/Mb for tTMB). Note that the correlation is higher when mutations are detected in both bTMB and tTMB (A). D) Patients who achieved a PR had median TMB higher than SD: 45 mut/Mb vs. 1 mut/Mb and PD: 45 mut/Mb vs. 5 mut/Mb. bTMB = blood-based tumor mutation burden; tTMB = tissue-based tumor mutation burden; PR = partial response; SD = stable disease; PD = progressive disease.

**Figure 2.** Mutational landscape of baseline plasma samples from 30 patients with detectable mutations. The most frequent mutations identified were: TP53 in 16 (53%), NOTCH2 and PKHD1, in respectively 5 (17%) and 5 (17%) patients. TMB = Tumor mutation burden; PR = Partial response; SD = Stable disease; PD = Progressive disease; yrs = years; IO = immunotherapy; RT = radiotherapy.

**Figure 3.** Change in VAF of ctDNA mutations encountered at baseline compared to cycle 2.
A) Anal squamous cell carcinoma; B) MSI-H endometrial carcinoma; C) Triple negative breast cancer. VAF = variant allele frequency; ctDNA = circulating tumor DNA; MSI-H = microsatellite instability-high; AF = allele frequency; C2 = cycle 2
Figure 1

A. Scatter plot showing the relationship between bTMB and tTMB with r = 0.71, p = 0.002. The plot distinguishes between concordant and discordant samples.

B. Scatter plot showing the difference in TMB against the difference in time (months) with best response RECIST categories indicated.

C. Scatter plot similar to A, showing r = 0.48, p = 0.02.

D. Scatter plot showing the best response RECIST categories with PD, PR, and SD represented by different symbols.
Figure 3

A

B

C

mutation

- BLM p.D44H
- FANCE p.E82Q
- PRSS3 p.D107H
- BTK p.L198F
- FANCN p.E1212Q
- PRSS3 p.E31D
- C11orf30 p.Q810*
- FAT1 p.L3767P
- RAD54L p.E122K
- CASP8 p.G339E
- FGF3 p.M631I
- RB1 p.E119Q
- CDKN1B p.S161C
- FGF3 p.S249C
- RB1 p.E894K
- CHD4 p.N1113S
- KIT p.G32A
- RUNX1T1 c.−23→1G>C
- CREBBP p.Q943E
- MECOM p.D212N
- SMAD3 p.R292T
- CTTNB1 p.L139F
- MECOM p.G259A
- SMARCA4 p.M949I
- CYLD p.K829N
- MPL p.Q18
- SMARCA4 p.Q201H
- DICER1 p.P934S
- MTOR p.E1871K
- SOX1 p.E103K
- DLL3 p.G223A
- PARP1 p.R878Q
- TGFB2 p.K1058fs*19
- EGFR p.T790K
- PBRM1 p.M1331I
- TPS3 p.R248Q
- EP300 p.Q965*
- PDGFRA c.1654→1G>C
- TP53 p.R1280T
- ERBB3 p.K736N
- PIK3CA p.E545K
- TSC2 p.Q35H
- ERBB3 p.S1083F
- PKHD1 p.D3139H
- TTF1 p.Q146*
- ERBB4 p.S1286F
- POLH p.S512*
- ZNF217 p.E914*

mutation

- APC p.E461Q
- LYN p.P79L
- ARID1A p.G324Afs*39
- MEF2B p.R127*
- ARID1A p.Q439H
- MGMT p.V186M
- ARID1A p.W1073*
- MSH2 p.L440P
- ATR p.S2485F
- NSD1 p.N1222Kfs*4
- BAX p.E41Rfs*19
- NSD1 p.R2039H
- BRD4 p.A1306V
- PALB2 p.M296*
- CDKN1A p.A39V
- PALLD p.G203V
- CHD4 p.F1174C
- PARP1 p.A483Qfs*3
- CHD4 p.R572*
- PIK3CA p.Q546H
- CHD4 p.R975H
- QKI p.K134Rfs*14
- EXT2 p.R557W
- RB1 p.R320*
- FGF4 p.R529Q
- SETD2 p.T305Qfs*35
- FLCN p.H429Tfs*39
- SPRY4 p.T221M
- GATA3 p.A146T
- TAP2 p.L86*
- GATA4 p.P253L
- TGFB2 p.K128Afs*3
- GSTT1 p.P189S
- TUBB4A p.R306C
- KMT2B p.L161Ffs*6
- YAP1 p.F96del

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