Changes in T-cell subsets and clonal repertoire during chemoinmunotherapy with pembrolizumab and paclitaxel or capecitabine for metastatic triple-negative breast cancer


ABSTRACT

Background Chemoimmunotherapy is a standard treatment for triple-negative breast cancer (TNBC), however, the impacts of different chemotherapies on T-cell populations, which could correlate with clinical activity, are not known. Quantifying T-cell populations with flow cytometry and T-cell receptor (TCR) immunosequencing may improve our understanding of how chemoimmunotherapy affects T-cell subsets, and to what extent clonal shifts occur during treatment. TCR immunosequencing of intratumoral T cells may facilitate the identification and monitoring of putatively tumour-reactive T-cell clones within the blood.

Methods Blood and tumor biopsies were collected from patients with metastatic TNBC enrolled in a phase Ib clinical trial of first or second-line pembrolizumab with paclitaxel or capecitabine. Using identical biospecimen processing protocols, blood samples from a cohort of patients treated for early-stage breast cancer were obtained for comparison. Treatment-related immunological changes in peripheral blood and intratumoral T cells were characterized using flow cytometry and TCR immunosequencing. Clonal proliferation rates of T cells were compared based on intratumoral enrichment.

Results When combined with pembrolizumab, paclitaxel and capecitabine resulted in similar peripheral T-cell subsets. Their effects were more modest than that observed following curative-intent dose-dense anthracycline and cyclophosphamide (ddAC) (average fold-change in CD3+ cells, capecitabine: −0.42, paclitaxel: −0.56, ddAC: −1.21). No differences in T-cell clonality or richness were observed following capecitabine or paclitaxel-based treatments. Regression modeling identified differences in the emergence of novel T-cell clones that were not detected at baseline (odds compared with ddAC, capecitabine: 0.292, paclitaxel: 0.652). Pembrolizumab with paclitaxel or capecitabine expanded T-cell clones within tumors; however, these clones did not always expand within the blood. Proliferation rates within the blood were similar between clones that were enriched and those that were not enriched within tumors.

Conclusion Chemoimmunotherapy for metastatic TNBC with pembrolizumab and capecitabine or paclitaxel resulted in similar peripheral T-cell subset lymphodepletion without altering T-cell clonal diversity. Regression modeling methods are applicable in immune monitoring studies, such as this to identify the odds of novel T-cell clones emerging during treatment, and proliferation rates of tumor-enriched T-cell clones.

INTRODUCTION

Recent phase III trials in metastatic triple-negative breast cancer (mTNBC) have demonstrated improved patient outcomes when chemotherapy was combined with anti-programmed death 1/ligand 1 (anti-PD-1/L1) monoclonal antibodies. The IMpassion130 study demonstrated improved progression-free survival (PFS) with nab-paclitaxel and atezolizumab (PFS anti-PD-L1, HR=0.62, p<0.001), and the Keynote-355 study demonstrated an improved PFS and overall survival (OS) with pembrolizumab plus various chemotherapies (nab-paclitaxel, paclitaxel, or gemcitabine with carboplatin) (PFS anti-PD-1, HR=0.65, p=0.0012). Despite these successes, not all chemoimmunotherapy regimens in all settings have improved outcomes for patients with metastatic disease. For example, neither first-line atezolizumab and paclitaxel nor later-line pembrolizumab and capecitabine improved OS or PFS, respectively. These discordant responses could be due to differential immune effects of the chemotherapy backbone.
The observation that the clinical activity of chemoimmunotherapy varies according to disease stage remains unexplained. For example, in stage II and III TNBC, pembrolizumab with chemotherapy improved outcomes across both PD-L1-negative and PD-L1-positive subgroups, whereas in metastatic disease the benefit of pembrolizumab with chemotherapy is restricted to the PD-L1-positive subgroup. For this reason, one aim of this biomarker study was to characterize the initial state of peripheral blood T-cell populations of mTNBC. T-cell subset quantities and clonal T-cell receptor (TCR) repertoire diversity were assessed in a phase Ib trial cohort of mTNBC patients, compared with a cohort of early-stage breast cancer (ESBC) patients. A second aim was to assess for changes to T-cell subset quantities and TCR clonal repertoire diversity in patients with mTNBC after combined pembrolizumab and capecitabine or paclitaxel. Effects of chemoimmunotherapy were compared with those of dose-dense anthracycline and cyclophosphamide (ddAC), which is a commonly used regimen that many patients with mTNBC receive prior to their diagnosis of metastatic disease; studying its effects on T cells enhances our understanding of the initial T-cell milieu of patients with mTNBC. Importantly, the samples from early-stage patients receiving ddAC were collected contemporaneously using identical biospecimen collection/processing methods and assays.

Measures of T-cell clonal diversity, such as richness (the numbers of unique T-cell clones) and clonality (evenness of clonal abundance within a population), have been correlated with outcomes following immune checkpoint antibody therapy in various cancer types (lung cancer, melanoma, pancreatic cancer, and urothelial cancer), as well as following chemotherapy in breast cancer. One explanation for this is that peripheral T cells are a reservoir of tumor-reactive T cells, thus, the richness or clonality of peripheral T cells may be useful as a surrogate measure for the ability to mount a successful antitumor response. However, one downside of generalized T-cell diversity metrics is that they cannot distinguish between clonal shifts that are attributable to specific antitumor activity from bystander activation or infection. Simultaneous detection of T-cell clones in both tumor and peripheral blood compartments using TCR immunosequencing has been demonstrated in a variety of cancer types. Proliferation of tumor-associated T-cell clones in the peripheral blood as a consequence of localized T-cell activation during treatment for mTNBC remains relatively unexplored. Our last aim was to determine whether intratumoral T-cell clones undergo greater proliferation, as measured by abundance, within the peripheral blood following therapy compared with T-cell clones restricted to the blood. If so, this would have represented a novel surrogate biomarker to measure treatment-related expansion of putative tumor-reactive T-cell clones. To measure the proliferation rate, we applied regression-based methods to interrogate the clonal dynamics of T-cell populations.

Monitoring peripheral blood T-cell subsets and T-cell clonal diversity during treatment can elucidate the variable biological effects of various chemoimmunotherapy regimens, particularly in the context of small investigator-initiated studies where samples can be uniformly collected and processed. In previous studies in melanoma, the percentage of individual peripheral T-cell subsets measured by flow cytometry was correlated with clinical response to anti-PD-1, which could be related to the fact that peripheral T cells can be an important source of tumor-reactive T cells. Additionally, overall lymphocyte count has correlated with clinical benefit from chemotherapy in breast cancer. Both paclitaxel and capecitabine are commonly used for the palliative treatment of mTNBC. Their immunological effects have been reported yet their effects when combined with checkpoint blockade have not been examined. We present the first comparison of the effects of pembrolizumab combined with paclitaxel or capecitabine on T-cell subset quantities and T-cell clonal repertoire diversity during first or second-line treatment of patients with mTNBC.

**METHODS**

**Participants and sample collection**

Participants with mTNBC (n=29) were enrolled in a phase Ib clinical trial between 2016 and 2018 at Providence Cancer Institute (Portland, OR) and Cedars-Sinai Medical Center (Los Angeles, California, USA) (NCT02734290). This study evaluated the safety of first or second-line pembrolizumab 200 mg intravenously on day 1 of a 3-week cycle, combined with either paclitaxel 80 mg/m² intravenously weekly on days 1, 8, 15 of a 3-week cycle (n=15) or oral capecitabine 2000 mg two times per day on days 1–7 of a 2-week cycle (n=14); chemotherapy was selected by the treating physician. All patients receiving paclitaxel additionally received dexamethasone 10–12 mg intravenously on cycle 1 day 1. Typically, this was continued weekly until day 15, when a taper to half the previous dose was attempted with subsequent discontinuation at cycle 2 by the treating clinician’s discretion. Only two of nine patients for whom these data are available continued to receive steroids beyond the third week of paclitaxel treatment. Peripheral blood samples were collected at baseline and every 3 weeks for the first 12 weeks of treatment. Tumor samples were collected at baseline and, when possible, a second biopsy was performed between weeks 5–8. For nine participants, the two biopsies were site matched. This trial met its safety endpoint of at least 6 weeks of treatment without dose discontinuation in 100% of patients in the capecitabine arm (14/14) and 87% (13/15) of patients in the paclitaxel arm. Response and safety data are reported elsewhere. Briefly, in the capecitabine arm, overall response rate (ORR) by response evaluation criteria in solid tumors V.1.1 was 43% (complete response (CR)=1, partial response (PR)=5, stable disease, SD=2, progressive disease (PD)=7) while...
in the paclitaxel arm, ORR was 29% (CR=2, PR=2, SD=1, PD=8) with one patient not evaluable at the cut-off but later assessed as a PR (table 1).

A contemporaneous cohort of patients with ESBC (n=29) was enrolled on a biospecimen collection protocol at Providence Cancer Institute. This presented a unique opportunity to characterize and then compare the immunological effects of curative-intent chemotherapy with chemoimmunotherapy using identical biospecimen protocols and assays. Nineteen subjects received four cycles of doxorubicin 60 mg/m² intravenously and cyclophosphamide 600 mg/m² on day 1 of a 14-day cycle, as part of the standard-of-care curative-intent therapy for stage II–III TNBC. To augment the sample size, patients with non-TNBC breast cancer subtypes were included in this analysis if they received the same dose and schedule of ddAC (table 1). Serial blood collections were performed during treatment at baseline and weeks 3, 5 and 9, corresponding to the doxorubicin-cyclophosphamide treatment phase, and when available an additional collection was performed between weeks 18–22. All patients provided written, informed consent.

Flow cytometry
Flow cytometry was used to determine CD45⁺CD3⁺ T-cell quantities and percentages at the Earle A. Chiles Research Institute (Portland, OR). Fresh whole blood was used to avoid deleterious effects of cryopreservation on cell count estimation. Blood samples collected from patients enrolled at Cedars-Sinai constituted the minority of specimens (n=8) and were collected using the Cyto-Chex BCT tube (Streck La Vista, NE), shipped to the Earle A. Chiles Research Institute, and analyzed within 24 hours of collection. In a separate validation cohort, absolute T-cell counts using this approach were highly correlated with counts obtained from heparinized tubes (data not

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CR, complete response; ddAC, dose-dense anthracycline and cyclophosphamide; H, trastuzumab; HER2, human epidermal growth factor receptor 2; HR, hormone receptor; IQR, interquartile range; P, pertuzumab; PD, progressive disease; PR, partial response; SD, stable disease; T, paclitaxel; TNBC, triple-negative breast cancer.
shown). The detailed protocol, including the gating strategy was described previously. The following CD3+ CD4+ and CD3+ CD8+ T-cell subsets were measured: naive (CD45RA+/CCR7+); central memory (CM) (CD45RA-/CCR7+); effector memory (EM) (CD45RA-/CCR7-); EM cell re-expressing CD45RA (EMRA) (CD45RA+/CCR7+). CD4+ T regulatory cells (Tregs) were characterized as CCR4+, CD127low, CD25high.

TCR immunosequencing

For assessment of T-cell clones, DNA was purified from PBMCs using the AllPrep DNA/RNA/miRNA Universal Kit (50) (QIAGEN, Cat # 80224) with QIAshredder Kit (50) (QIAGEN, Cat # 79654) for homogenization. DNA was purified from formalin-fixed paraffin-embedded (FFPE) tumor specimens using the AllPrep FFPE Kit (50) (QIAGEN, Cat # 80234). Purified DNA was submitted for high-throughput deep sequencing using the immunoSEQ TCRB Assay (Adaptive Biotechnologies, Seattle, WA). mTNBC samples collected at baseline, weeks 4, 7 and 13 and ESBC samples collected at baseline, weeks 3, 9, and between weeks 18 and 22 were submitted.

Analysis of TCR immunosequencing data

Various statistical methods were employed to evaluate T-cell clones over time in the peripheral blood and tumor. The T-cell richness, clonality, and differential abundance metrics are commonly cited in the literature and can be directly accessed using the immunoSEQ Analyzer software platform (Adaptive Biotechnologies, Seattle, WA), and are summarized below. We applied regression methods (not yet cited in the literature for this application) to evaluate T-cell clones over time, and to compare effects across treatment groups.

T cell richness

Clonal richness is one facet of overall diversity of T-cell clones within a population and describes the total number of unique clones in a sample (identified by each clone’s CDR3β amino acid sequence). Because cell counts across different blood draws are not standardized, and because the overall sample size can influence the number of unique clones, we present two separate metrics, a unique clone count (which is unadjusted for sample size), as well as T-cell richness, which is adjusted for sample size. T-cell richness is calculated as the average number of unique T-cell clones across 10 random down-sampling events (downsampled to 5620 templates for peripheral blood samples, and 248 templates for tumor biopsy samples). The number of templates represent the number of input T cells after a bias-corrected normalization of TCR immunosequencing reads.

Simpson’s clonality

Clonality is another facet of diversity which describes the evenness of the population. Here, it describes whether T cells within a population are uniform (all the same clone, monoclonal) or diverse (many different clones, polyclonal). It is calculated as the square root of the Simpson’s diversity index, which incorporates the number of species present and their abundance. High scores indicate an uneven mono/oligo-clonal T-cell population, whereas low scores indicate a more even, polyclonal T-cell population.

Differential abundance

To assess whether certain T-cell clones are expanded/contracted in one sample vs another, a statistical test must account for differences related to overall sample size. For this, we utilized Fisher’s exact testing with a beta-binomial model significance test and multiple hypothesis correction by the Benjamini-Hochberg method. We also used this method to identify tumor-enriched clones by comparing tumor vs blood samples collected at the baseline timepoint.

Longitudinal analyses

Serial trends in T-cell subtype or clonal concentrations across multiple time points were analyzed using R software, V.4.0.3, and the R package ‘limma’. Various regression methods were employed, as summarized below.

Longitudinal cellular abundance analysis

To assess the overall rate of T-cell subset expansion or contraction following treatment, we applied a mixed effects longitudinal regression model (with time being a fixed effect, and cell count being the endpoint). This provides an estimate of the rate of change over time (and 95% CI) for the cellular subset of interest. Variations related to study subject are accounted for by including study subject as a random effect in the model.

Longitudinal T cell clone detection analysis

Because chemotherapy is lymphodepleting, we developed linear mixed effects models to quantify relative differences in T-cell clonal loss related to therapy. These models have been used in the longitudinal study of CD4+ T cell counts and HIV DNA abundance during antiretroviral treatment for HIV, though to our knowledge have not been applied previously to TCR immunosequencing data. We used a logistic approach to estimate of the odds (ranging from 0 to 1) of a T cell clone would become undetectable during treatment, with values close to 0 indicating that the therapy results in loss of all T-cell clones, and values close to 1 indicating that all T-cell clones remain detectable over time. For the model, T cell detectability is a binary variable, time is a fixed effect and the subject is a random effect.

A similar strategy was employed to evaluate the OR of the emergence/detection of T-cell clones that were undetectable before starting treatment relative to a reference treatment. The model has a similar structure to the above logistic regression model, but it estimates the odds (ranging from 0 to 1) of detection of the subgroup of clones that were not detected in the first sample collection, but were detected later relative to a reference treatment. Values close to 0 would indicate that new T-cell clones are less likely to be detected compared with the
Longitudinal clonotype abundance analysis
We estimated the mean T-cell clonal proliferation rate related to each treatment using regression models with similar variables as described above. Because there is no reported gold standard method, we explored various regression methods, and found similar outcomes. Details of the various models, including assumptions, merits, and caveats, are summarized in online Supplemental Table 1. In the results we present outcomes of a Poisson generalized linear mixed model, applied to the subgroups of clones which were enriched within the tumor or were neither enriched within the tumor or not detected in the tumor at all.

We provide a visual method for illustrating the spectrum of T-cell clonal proliferation rates across different subgroups. We calculated the proliferation rate based on the abundance of individual T-cell clones per week, and plotted them in rank order from lowest to highest (by their percentile ranking). To account for possible variations in the rate of clonal proliferation attributable to the starting prevalence, we defined a subgroup of tumor-absent clones with similar baseline frequency to the tumor-present clones. Because these data may be influenced by the overall size of the sample, to compare the subgroups, we randomly sampled 100,000 T-cell clones for each group, with replacement as necessary.

All other statistical analyses to compare T-cell subset quantities and percentages, and diversity metrics were performed using R software, and included two-sample t-test, Kruskal-Wallis, analysis of variance, and Pearson’s linear correlation coefficient testing as indicated. P values are provided to aid in hypothesis generation but are not corrected for multiple comparisons.

RESULTS
Baseline T-cell profile of mTNBC patients
We first characterized baseline peripheral T-cell populations of ESBC and mTNBC patients by flow cytometry and TCR immunosequencing. There were no appreciable differences in T-cell subset sizes within the contemporaneous ESBC cohort according to histology (online supplemental figures 1 and 2). Because there were no differences, data from the entire cohort of ESBC were combined. Patients with mTNBC had fewer lymphocytes than patients with ESBC (CD45+ ESBC: 2.2×10⁹ cells/L, vs mTNBC: 1.2×10⁹ cells/L, p<0.001) (figure 1A). This relation was observed across all T-cell subsets, including CD3+ cells, CD4+ cells, CD8+ cells, and naïve, CM, EMRA, and regulatory T cells (figure 1B-E, online supplemental figure 1). Figure 1E lists the average ratio of cell counts across the various T-cell subsets, comparing mTNBC to ESBC. Despite reduced T-cell subset quantities among the mTNBC cohort, the percentages of T-cell subsets relative to all T cells were similar in the mTNBC vs ESBC cohorts (online supplemental figure 2).

Using TCR immunosequencing, we evaluated T-cell clonal diversity in the mTNBC vs ESBC cohorts. The two cohorts exhibited similar mean T-cell richness (ESBC: 4626 unique rearrangements per 5260 TCR molecules, mTNBC: 4440 unique rearrangements per 5260 TCR molecules, p=0.40, t-test) and clonality (ESBC: 0.055, mTNBC: 0.055, p=0.98, t-test) (figure 2A,B). There were no differences in T-cell richness or clonality between the histological subgroups among the ESBC cohort (online supplemental figure 3). We evaluated the relative distribution of T-cell clones according to their abundance and found similar proportions of high-frequency clones between the cohorts, but differences in lower and moderate-frequency clones (figure 2E).

Because chemotherapy can be associated with long-term alterations to peripheral T-cell profiles, we compared different subgroups of mTNBC participants: chemotherapynaïve (n=5), most recent chemotherapy >12 months ago (n=16), and most recent chemotherapy 6–12 months ago (n=8). The average baseline CD3+ T-cell count did not vary significantly among these groups, except for CD8+ EM cells (online supplemental figures 1 and 2). Comparisons of richness and clonality were limited due to lower numbers of chemotherapynaïve patients with available TCR immunosequencing data (data not shown).

Effects of chemoimmunotherapy on peripheral T-cell subsets and TCR repertoire in mTNBC
We observed similar declines in CD3+ T cells over time whether patients received pembrolizumab with capecitabine or paclitaxel (average CD3+ T-cell fold change capecitabine: −0.42, paclitaxel: −0.56, p=0.80, t-test). By comparison, patients in the ESBC cohort treated with ddAC experienced a greater loss of CD3+ T cells (average fold change: −1.21). Similarly, the rate of T cell loss across all major T-cell subsets, including Tregs, was greater after ddAC than either of the pembrolizumab-based chemotherapy regimens (figure 3, online supplemental table 2). There were no significant changes to TCR repertoire richness or clonality with either capecitabine or paclitaxel combined with pembrolizumab, whereas ddAC resulted in reduced T cell richness but no change in clonality (figure 4A,B). There were not differences in the number of significantly expanding (figure 4C) or contracting clones (figure 4D) between patients treated with pembrolizumab combined with capecitabine or paclitaxel. Following ddAC there were more expanded clones and fewer contracted clones than observed after either pembrolizumab-based regimen. However, this finding did not reach statistical significance (figure 4C,D). The expanding clones identified during each treatment were durable and detectable at later time periods (capecitabine: 85%, paclitaxel: 85%; ddAC: 80%). In summary, using these generalized metrics, we found minimal differences in T-cell clonality or diversity between the two
pembrolizumab-based regimens, whereas the curative-intent ddAC regimen was associated with a loss of overall diversity and evidence of more T-cell clonal expansions.

We applied regression methods to interrogate further the effect of these therapies on the T-cell clonal repertoire. Regarding T-cell clones, we confirmed that the rate of clonal loss between the pembrolizumab-based arms were similar, and lower than observed after ddAC (figure 4E). On the other hand, pembrolizumab with chemotherapy resulted in fewer T-cell clones emerging during treatment than arose during ddAC (figure 4E). In summary, these data may suggest that ddAC administered for early-stage treatment facilitates robust peripheral T-cell clonal proliferation despite great lymphotoxicity.

**Changes to intratumoral T cells during chemoimmunotherapy for mTNBC**

From the mTNBC patients, we obtained nine pairs of pretreatment and on-treatment tumor samples from the same lesion, providing an opportunity to evaluate changes to intratumoral lymphocytes during treatment. Overall, changes in intratumoral T-cell richness and clonality varied across patients and no specific trend could be detected (figure 5A,B). However, pretreatment/post-treatment comparisons were not evaluable for all pairs because of low numbers of TCR templates in some samples. The number of T cells detected by TCR immunosequencing increased in some pairings, but not in all (figure 5C); however, the fraction of T cells among all nucleated cells increased in most samples (figure 5D).

On average, each patient had 771 (range 1–4964) T-cell clones that were present in both baseline and on-treatment biopsies. On average, these clones comprised 19% of the baseline clones (range: 0.1%–36%) and 17% of the on-treatment clones (range: 3%–41%). In five subjects we identified T-cell clones that significantly expanded or contracted following therapy at an alpha level of 0.05 (figure 5F). Expanding T-cell clones were more likely to be detected in the peripheral blood, when compared with those that contracted (expanding: 80%; contracting: 43%). However, expansion or contraction of T cell clones...
in the tumor did not always correlate with expansion or contraction in the peripheral blood (figure 5E). Out of the three patients with expansion of at least one clone in both the tumor and the blood, only one patient had a PFS greater than the average for the entire cohort of mTNBC patients (patient PFS: 258 days; cohort mean PFS: 146.3 days, range: 21–426 days) (figure 5E). TCR CDR3β amino acid sequences of differentially-abundant clones within tumors (online supplemental table 3) did not match any previously-identified sequences within publicly available V(D)J databases McPAS-TCR⁴⁹ and VDJ-db⁵⁰ though the amount of available data is currently limited.

**Figure 2** Baseline TCR repertoire diversity metrics for early-stage breast cancer and metastatic triple-negative breast cancer cohorts as measured by (A) richness, (B) clonality, (C) total TCR templates per sample and (D) total unique clones. Significance testing is by the two-sample t-test. (E) Average cumulative abundance of clones grouped by prevalence. TCR, T-cell receptor.

**Use of tumor specimens to identify and track tumor-enriched TCR clones**

We explored whether longitudinal trends of peripheral T-cell clone abundance differed based on the clone’s presence within pretreatment tumor. Pretreatment tumor and serial blood samples were available from 19 mTNBC clinical trial patients which facilitated the identification of intratumoral T-cell clones, some of which are putatively tumor-reactive. Interestingly, there were seven clones which were expressed in both tumor and blood compartments, and detected in more than one person. Expression of a shared clone was not necessarily associated with clinical response (online supplemental table 4). Among clones present within the tumor, their abundance was loosely correlated to their average relative abundance in the blood (figure 6A), raising the alternate hypothesis that T-cell clones in the tumor are related to tissue perfusion by peripheral T-cell clones, which are not necessarily tumor-reactive.

To explore these contrasting possibilities, we looked for differences in trends of clonal abundance among a subset of intratumoral T-cell clones that were enriched within the tumor, relative to the peripheral blood (using the differential abundance statistical test). We identified 1914 tumor-enriched clones among the 19 patients (average per patient: 101, range: 0–469) out of 145,534 tumor-present T-cell clones (average per patient: 7660, range: 79–54,714) (online supplemental figure 4B). On average, tumor-enriched clones comprised 0.79% of the total detected TCR sequences within the peripheral blood compartment and 20.2% within the tumor compartment (online supplemental figure 4B). Tumor-present
clones comprising the remaining 79.8% of tumor TCR sequences comprised 18.7% of TCR sequences within the peripheral blood. There were no tumor-enriched clones from baseline samples which were shared between subjects.

Tumor-present and tumor-enriched clones were more abundant in the blood on average than tumor-null clones (average TCR molecules per clone at week 7, tumor-enriched: 8.8, tumor-present: 1.8, tumor-absent: 0.39), but the average rate of change in the blood was similar across these three groups (figure 6B, online supplemental table 5). The quantile plot (figure 6C) illustrates the individual slopes of clone abundances (rate of change) identified in the groups: tumor-enriched, tumor-non-enriched, and peripheral blood matched and unmatched for starting abundance. As illustrated, the

![Figure 3](image-url)
abundance in the blood of most T-cell clones is stable over time across groups, and rare clones expand or contract.

**DISCUSSION**

In this post hoc analysis of potential biomarkers, we employed comprehensive T-cell monitoring using fresh whole-blood flow cytometry and blood/tissue TCR immunosequencing of blood and tissue to provide insights into the starting T-cell profile of patients with mTNBC, and the subsequent effects of palliative chemoimmunotherapy on T-cell populations. We examined alterations in T-cell populations during ddAC treatment in an early-stage cohort as a comparator to aid hypothesis generation regarding the development of immunosuppression which may be prevalent in mTNBC patients. We addressed three aims: (1) to characterize baseline T-cell profiles of
patients with mTNBC by T cell subset quantity and clonal repertoire; (2) to characterize the effects of two different chemoimmunotherapy regimens on T-cell subsets and clonal repertoire; and (3) to explore whether the peripheral proliferation rate of clones that are enriched within the tumor is different than that of non-enriched clones.

We characterized the baseline peripheral blood T-cell counts and clonal diversity in patients with mTNBC relative to ESBC because baseline differences in the T-cell profile may contribute to observed differences in clinical efficacy of chemoimmunotherapy between early and metastatic stage TNBC. In our analysis, mTNBC patients exhibited global reductions in peripheral T-cell quantities relative to ESBC counterparts, with absolute cell counts of T-cell subsets in the metastatic setting ranging from 36% to 80% of counts observed in ESBC. Furthermore, while mean T-cell clonal diversity scores were similar in mTNBC and ESBC, there was greater variability in these metrics within the mTNBC cohort, highlighting the heterogeneity of patients within the metastatic cohort.
We also detected differences in the cumulative proportion of low/mode-abundance T-cell clones in the overall TCR repertoire. Overall, these data suggest that the systemic T-cell milieu is suppressed in patients with mTNBC. Whether this directly relates to differences in observed efficacy of checkpoint inhibition in metastatic and early-stage TNBC is worthy of further study.

Important questions remain regarding whether our observation of relative lymphodepletion in patients with mTNBC is a consequence of previous therapies resulting in bone marrow suppression, and whether T cells in patients previously treated with chemotherapy have reduced functional capabilities. Although the dataset is too small to provide definitive conclusions, we observed that the timing of recent curative-intent therapy in mTNBC patients did not impact T cell concentrations or richness. There were some numerical differences between the chemotherapy-naïve patients, and those previously treated with chemotherapy, which is supported by literature that curative-intent chemotherapy results in long-term T cell depletion, particularly of CD4+ subsets. Understanding the long-term effects of previous therapies on T-cells is worthwhile as chemoimmunotherapy appeared to confer greater benefit in PFS for chemotherapy-naïve patients in the IMpassion130 and Keynote-355 trials. Additionally, in a previous phase II study, pembrolizumab failed to improve PFS when combined with capecitabine in later lines of therapy, which contrasts with our phase Ib trial that reported a more favorable PFS of pembrolizumab plus capecitabine when administered earlier in the disease course. Other factors, such as tumor burden and age may contribute to immunosuppression and may be relevant in patients with mTNBC.

We characterized the effect of pembrolizumab plus capecitabine or paclitaxel on T-cell subsets and the TCR clonal repertoire. Both chemotherapies are used
commonly in the palliative treatment of mTNBC and are safe in combination with pembrolizumab. While immune effects of these chemotherapies as single agents have been described, the effect of adding pembrolizumab has not been examined. The effects could be beneficial or harmful. Capcitabine and its active moiety, 5-fluorouracil, may enhance peptide presentation by MHC-I in tumor cells, deplete myeloid-derived suppressor cells (MDSCs), and increase T-helper 17 cells (T₇₁₇). Paclitaxel repolarizes macrophages to an antitumor M1 phenotype, depletes MDSCs, and promotes dendritic cell maturation and function. There is additional concern that corticosteroid prophylaxis, routine with paclitaxel therapy, but not capcitabine, may contribute to suppressed T cell viability, signaling, and function. When capcitabine and paclitaxel were combined with pembrolizumab, we did not observe meaningful differences in T-cell subset losses using conventional surface markers nor in T-cell clonal expansion using previously established methods of assessing T-cell clonal diversity or differential abundance. Using a novel application of logistic regression, emergence of novel clones was more likely to occur following paclitaxel than capcitabine chemoimmunotherapy. Emergence of new clones was even more likely during ddAC. This finding requires additional confirmatory study, but generates interest because novel clones which emerge during treatment may have the potential to target tumor-associated neoantigens. It is possible that ddAC stimulated the expansion of more clones due to doxorubicin’s immunogenic effects, or that the more severe lymphopenia induced by ddAC triggered rapid endogenous proliferation of T cells. Additional evaluations to determine any differences between chemotherapy backbones on immune cell populations could include the quantification of additional immune cell subsets, characterization of T-cell function using direct T-cell functional assays or single-cell RNA sequencing or proteomic assays.

The third goal was to characterize dynamics of intratumoral versus peripheral T-cell clonal expansion, in hopes that we could identify a peripheral blood surrogate biomarker for tumor-specific T-cell response related to chemoimmunotherapy. TCR sequencing of tumor digests did not reveal a pattern of changes in intratumoral T-cell clonal diversity related to treatment; however, it identified statistically significant T-cell clonal expansion events in site-matched specimens. When tracked individually, T-cell clones enriched within the tumor (including those found to be expanding) were often detected within the blood, but did not necessarily expand within the blood. Among each population of T-cell clones there is a range of proliferative rates within the blood, with the vast majority of clones undergoing no proliferation and rare outliers either proliferating or contracting. In summary, these data suggest that expansions of intratumoral T-cell clones are detectable within the blood, but these are rare even when focusing on T-cell clones enriched within the tumor, and clones that expand within the tumor can easily be overlooked if one relies solely on the blood to identify potentially tumor-reactive T-cell clones. In future experiments, TCR sequencing of T-cell subsets that are more likely to be tumor-reactive (such as CD103/CD39 double-positive T cells) or T cells with CXCL13, CD39, TOX, and PD-1 expression could more accurately identify tumor-reactive T-cell clones.

There are limitations to this biomarker analysis. First, as a biomarker-focused investigator-initiated trial, the sample size is small relative to phase III trials such as Keynote-355 and Impassion130, and therefore, the findings are exploratory and not powered to detect small differences across the treatment arms. However, one virtue of this analysis was that the specimens were processed and analyzed in one laboratory using a standardized protocol and non-cryopreserved blood. This type of analysis is often impossible to conduct using specimens from large, multiinstitutional clinical trials. Limitations regarding the comparison of treatment effects arise because patients were not randomized between capcitabine and paclitaxel, and the contemporaneous EBSC cohort comprised a variety of tumor histotypes receiving ddAC without the inclusion of pembrolizumab. Therefore, it is possible that some of the differences attributed to ddAC are in fact related to the omission of pembrolizumab, differences in tumor subtype, or the immune contexture of early-stage disease. Therefore, it is important that these findings are confirmed in a more homogenous dataset. We will later report immunological profiles of a larger cohort of n=30 stage II/III TNBC patients receiving ddAC with pembrolizumab in an ongoing phase II investigator-initiated trial (NCT04373031). Meanwhile, our reported data offers useful insights for other investigators planning to study changes to T-cell populations during immunotherapy trials. A third limitation is that our analysis was restricted to whole-blood flow cytometry and whole-tumor TCR sequencing. For future studies, we advocate for single-cell multiomic characterizations of T-cells to capture TCR clonotype and functional state to better characterize T-cell activity and clonal proliferation in response to treatment.

CONCLUSION

We present longitudinal immunological profiling of the effect of pembrolizumab with capcitabine or paclitaxel in mTNBC. At baseline, patients with mTNBC, the majority of whom were pretreated, exhibited depletion of T-cell subsets and reduced T-cell clonal richness compared with untreated EBSC patients, but percent-ages of T-cell subsets and clonality were not significantly different between cohorts. We did not detect significant differences in T-cell subset losses when comparing patients treated with either of the two chemoimmuno-therapy arms. Exploratory application of logistic regression techniques revealed differences in the likelihood of emergence of novel clones between the various treatments. Changes to TIL populations were heterogeneous within nine paired tissue samples. Lastly, regression-based
statistical methods can characterize the peripheral proliferation rate of T-cell clones and are applicable in immune monitoring studies.

Author affiliations
1Earle A Chiles Research Institute, Providence Cancer Institute, Portland, Oregon, USA
2Knight Cancer Institute, Oregon Health & Science University, Portland, Oregon, USA
3Medical Data Research Center, Providence St Joseph Health, Portland, Oregon, USA
4Computational Immuno-Oncology and Bioinformatics Core, Earle A. Chiles Research Institute, Portland, Oregon, USA
5Adaptive Biotechnologies Corp, Seattle, Washington, USA
6Department of Medicine, University of Texas Southwestern Medical Center, Dallas, Texas, USA

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