Multimodality analysis confers a prognostic benefit of a T-cell infiltrated tumor microenvironment and peripheral immune status in patients with melanoma

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ABSTRACT

Background We previously reported results from a phase 1 study testing intratumoral recombinant poliovirus, lerapolturev, in 12 melanoma patients. All 12 patients received anti-PD-1 systemic therapy before lerapolturev, and 11 of these 12 patients also received anti-PD-1 after lerapolturev. In preclinical models lerapolturev induces intratumoral innate inflammation that engages antitumor T cells. In the current study, lerapolturev and postlerapolturev tumor biopsies and blood were evaluated for biomarkers of response.

Methods The following analyses were performed on tumor tissue (n=11): (1) flow cytometric assessment of immune cell density, (2) NanoString Digital Spatial profiling of protein and the transcriptome, and (3) bulk RNA sequencing. Immune cell phenotypes and responsiveness to in vitro stimulation, including in vitro lerapolturev challenge, were measured in peripheral blood (n=12).

Results Three patients who received anti-PD-1 therapy within 30 days of lerapolturev have a current median progression-free survival (PFS) of 2.3 years and had higher CD8+ T cell infiltrates in prelerapolturev tumor biopsies relative to that of 7 patients with median PFS of 1.6 months and lower CD8+ T cell infiltrates in prelerapolturev tumor biopsies. In peripheral blood, four patients with PFS 2.3 years (including three that received anti-PD-1 therapy within 30 days before lerapolturev and had higher pretreatment tumor CD8+ T cell infiltrates) had significantly higher effector memory (CD8+, CCR7-, CD45RA-) but lower CD8+PD-1+ and CD4+PD-1+ cells compared with eight patients with median PFS 1.6 months. In addition, pretreatment blood from the four patients with median PFS 2.3 years had more potent antiviral responses to in vitro lerapolturev challenge compared with eight patients with median PFS 1.6 months.

Conclusion An inflamed pretreatment tumor microenvironment, possibly induced by prior anti-PD-1 therapy and a proficient peripheral blood pretreatment innate immune response (antiviral/interferon signaling) to lerapolturev was associated with long term PFS after intratumoral lerapolturev in a small cohort of patients.

These findings imply a link between intratumoral T cell inflammation and peripheral immune function.

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INTRODUCTION

Both host-mediated antitumor immune responses and therapeutic strategies designed to reinvigorate host immune responses are subject to multiple tumor-driven immune suppressive mechanisms and protumorigenic signaling that occur in the tumor microenvironment (TME). Cellular and molecular pathway evaluation have shown that the TME can regulate tumor progression and that altering the TME can correlate with therapeutic benefits. In particular, pretreatment melanoma tumors with higher intratumoral CD8+ T cells and increased IFN-γ signaling within the TME, so-called ‘hot tumors’, have been associated with response to anti-programmed cell death protein 1 (PD-1) therapy. Anti-PD-1 therapy reinvigorates intratumoral inflammation by expanding stem-like effector T cell populations in the TME, thereby altering the composition within the TME. Alternatively, ‘cold’ tumors are characterized by minimal or absent T cell infiltration, and are less responsive to anti-PD-1 therapy. Among ‘cold’ tumors, phenotypes can include a lack of any appreciable T cell infiltrate or the peritumoral accumulation of T cells which are unable to enter the tumor, likely as a result of immunosuppressive myeloid cells or stroma in the TME. In melanoma, anti-PD-1 therapies have become the cornerstone of treatment for patients with metastatic disease and yet the majority of patients fail to respond; treatment failure...
is thought to be related in part to a ‘cold’ tumor phenotype and failure to remodel the TME.11 One approach for melanoma patients lacking responses to anti-PD-1 therapy is intratumoral delivery of agents designed to promote inflammation in the TME, which could potentially promote a ‘hot’ tumor phenotype.15–17 Indeed, lerapolturev treatment of ex vivo patient tumor slice cultures culminated in robust induction of PD-L1 expression on myeloid cells; intratumor therapy of murine melanoma tumors resulted in increased tumor infiltration of PD-L1-expressing CD4 and CD8 T cells and elicited functional antitumor CD8 T cell responses.16 17 These studies indicated potential for lerapolturev to sensitize melanoma tumors to anti-PD-1 therapy. Recently, we reported results of a phase 1 trial of intratumoral PVSRIPO in 12 patients with unresectable melanoma.18 In the phase 1 trial, four patients achieved an objective response while eight patients had no response; the injections were well tolerated. To determine immunologic effects in the TME and in the periphery, we performed tumor biopsies and blood analysis before and after lerapolturev. Here, we describe evaluation of the TME and peripheral blood analysis from melanoma patients receiving lerapolturev including the relationship of the pretreatment TME with clinical outcome, differences in pretreatment versus post-treatment tumor, comparison of results among the independent techniques used to study the tumor, and the relationship between innate activation potential in the peripheral blood and clinical outcome.

METHODS

All work was performed using Institutional Review Board approved protocols and patients signed written consent to participate. Trial details were published previously.18 Three patients received 1 injection of lerapolturev (patients 1–3), 3 patients received 2 total injections, each injection into a separate tumor (patients 4–6), 3 patients received 3 total injections each injection into a separate tumor (patients 7–9), and 3 patients received 3 total injections into the same tumor (patients 10–12).18 Tumor and peripheral blood were collected at multiple time points before and after lerapolturev. Pretreatment biopsies were collected 7–10 days prior to lerapolturev and post-treatment biopsies were performed 10 days after lerapolturev (for patients 1–9) and at median 30 days after last lerapolturev injection (for patients 11 and 12). In some cases, non-lerapolturev treated tumor was also biopsied. Flow cytometry analysis of tumor was conducted in real-time, shortly after sample collection with minimal processing to prevent exclusion or death of innate immune cells (eg, neutrophils, basophils, and eosinophils). A separate portion of each tumor was also preserved in paraffin. For each tumor specimen, a pathologist confirmed presence of viable tumor on H&E stains; if no viable tumor was noted, the sample was not used for analysis. Flow cytometry and in vitro stimulation assays were performed using banked frozen peripheral blood mononuclear cells (PBMCs) after isolation from blood.

Flow cytometry

Tumor tissue was collected and stored in MACS tissue storage solution at 4°C (Miltenyi). Storage time was 2–16 hours post-tumor collection. Tumor cells and tumor infiltrating immune cells were analyzed after tumors were processed using the Tumor Dissociation Kit and Gentle MACS mechanical dissociator (Miltenyi) following the manufacturer’s recommendations. The suspensions were then filtered three times through a 70 µm filter to remove undigested tissue and debris. Single cell suspensions were immediately incubated with the DuraClone IM Basic panel, which contains CD16-FITC, CD56-PE, CD19-ECD, CD14-PC7, CD4-APC, CD8-AF700, CD3-AF750, and CD45-KO (Krome Orange) as previously described, to identify immune cells in tissue.19 In addition, propidium iodide (Sigma-Aldrich, St. Louis, MO) was added as a live-dead marker to single cell suspensions. Cells were incubated for 15–20 min in the dark and then washed twice in phosphate-buffered saline (PBS) prior to data acquisition. Samples were analyzed on a 13-color CytoFlex flow cytometer (Beckman Coulter). Data were analyzed using Kaluza Software (Beckman Coulter). Comparisons were made between two groups (responders and non-responders) using unpaired t-tests.

For analyses of PBMCs, frozen PBMCs were thawed in AIM-V media (Gibco) containing DNase I (1 µg/mL; Roche) and incubated for 30 min at 37°C. Cells were pelleted and resuspended in PBS (Gibco) containing Zombie-NIR stain (1:500; BioLegend) for 30 min. The cells were washed in PBS containing 2% fetal bovine serum (FBS) (Sigma-Aldrich, St. Louis, MO) was added as a live-dead marker to single cell suspensions. Cells were incubated for 15–20 min in the dark and then washed twice in phosphate-buffered saline (PBS) prior to data acquisition. The following antibodies were added for staining at a 1:100 dilution: CD3-BUV737, CD8-BUV395 (from BD Biosciences); or CD4-BV510, CD25-PERCPc55.5, CD27-BV421, CD40L-PECF594, CD62L-FITC, PD1-BV711, CD127-PEcy5, CD45RA-APC, CD45RO-BV786, CD28-PECy7, CD56-BV605, and CCR7-PE (all BioLegend). After 1 hour cells were washed in PBS+2% FBS containing human Fc block (1:100, BioLegend). The following antibodies were added for staining at a 1:100 dilution: CD3-BUV737, CD8-BUV395 (from BD Biosciences); or CD4-BV510, CD25-PERCPc55.5, CD27-BV421, CD40L-PECF594, CD62L-FITC, PD1-BV711, CD127-PEcy5, CD45RA-APC, CD45RO-BV786, CD28-PECy7, CD56-BV605, and CCR7-PE (all BioLegend). After 1 hour cells were washed in PBS+2% FBS and analyzed on a Fortessa X20 alongside fluorescence minus one controls for CD28, CD27, and PD1. Gating and analyses were performed using FlowJo V.10 (BD Biosciences).

In vitro PBMC stimulation assay

For in vitro stimulation of PBMCs, laboratory grade PVSRIPO was generated in HeLa cells (American Type Culture Collection, ATCC) as previously described.20 Frozen, pretreatment PBMCs were thawed in AIM-V media (Gibco), treated with DNaseI (1 µg/mL; Roche)
for 30 min, counted using a Countess II automated cell counter (Thermo-Fisher), and plated in 6 wells of a 24 well plate with PBMCs at a density of 5×10⁵ cells per well in RPMI-1640 (Gibco) containing 10% FBS (Sigma-Aldrich) with no other additives or added cytokines. The following stimulants were added to separate wells: mock (vehicle); PVSRIPO (1×10⁷ plaque forming units (pfu)); lipopolysaccharide (LPS) (100 ng/mL; Invivogen); Pam3CSK4 (1 µg/mL; Invivogen); R848 (1 nM; Invivogen); or combined anti-CD3 (1 µg/mL clone OKT3; BioLegend), anti-CD28 (1 µg/mL clone CD28.2; BioLegend), and goat anti-mouse IgG (4 µg/mL; Jackson Immunoresearch). Cultures were incubated for 24 hours in a 37°C and 5% CO₂ incubator, after which supernatant was harvested. Supernatant cytokines were measured using the Human Anti-Virus Response LegendPlex panel (BioLegend) on a Fortessa X20 (BD Biosciences), and the mean fluorescence intensity for each analyte was determined using the manufacturer’s LegendPlex data analysis software (BioLegend). Fold mock control values were determined for each PBMC sample for each analyte.

**NanoString digital spatial profiling**

Formalin-fixed, paraffin-embedded (FFPE) tissue sections were analyzed using the NanoString GeoMx digital spatial profiling (DSP), a non-destructive method for high-plex spatial profiling of proteins and protein-coding genes. Detailed methods have been described. In brief, this method uses small photocleavable oligonucleotide ‘barcodes’ (PC oligos) covalently attached to in situ affinity reagents (antibodies and RNA probes) to provide unlimited multiplexing capability. The photocleavage light is projected onto the tissue slice using two-digital micromirror devices, containing 1 million semiconductor-based micromirrors allowing complete flexibility in the pattern of light utilized for high-plex digital profiling of the tissue. These spatial light patterns can be automatically configured to profile manually selected regions or molecularly defined compartments. Cell subset analysis can be done on a region with as low as 30 cells expressing the marker of interest. After a pathologist confirmed the presence of tumor on FFPE H&E stains, unstained slides were fluorescently stained to visualize the morphology for regions of interest (ROIs) selection and eventual proteomic or whole transcriptome analysis (WTA).

**Statistical analysis of NanoString DSP**

The protein panel included 59 proteins including internal-spike controls which were analyzed in selected ROIs. The raw data were normalized with a panel of three housekeeping proteins and three negative controls. The final step to control for variability in area and/or cellularity relies on determining the signal to noise ratio (SNR) which is calculated as the ratio of each probe over geometric mean of negative controls (mouse IgG₁, mouse IG2a and rabbit IgG). The SNR was then used to compare protein expression between our proposed ROI. The SNR used for comparisons for each sample represents the medians from 3 to 4 ROIs (tumor or cell type). The normalized DSP SNR data of protein expression were used to create clustered heatmaps. Each heatmap row represents a patient sample and each column corresponds to a protein, with the color and intensity of the boxes indicating relative expression. The heatmaps were created using the Broad Institute’s Morpheus software. Hierarchical clustering analysis was then performed on the heatmaps. The distance measure used was the 1—Spearman correlation coefficient. The average linkage method was used to determine the proximity between two clusters. These parameters enabled hierarchical clustering on both the rows and columns.

Similarly, the WTA analysis including 14442 human transcripts were analyzed from ROIs, normalized and the normalization values were used to make comparisons. To account and control for technical variability, a common set of external RNA controls developed by the External RNA Controls Consortium (ERCC) was used. In addition to the raw data being ERCC normalized, the raw data were normalized with a panel of three housekeeping proteins and three negative controls. After ERCC normalization a final step to control for variability in area and/or cellularity relies on determining the SNR, which is calculated as the ratio of each probe over geometric mean of negative controls. The SNR was then used to compare expression between our proposed ROI. The NanoString WTA values were analyzed using the t-test to compare the average ROIs of two groups and using a volcano plot to represent the differential expression of transcripts; fold change was calculated as mean group 1—means group 2 divided by mean group 2. If the p<0.05 and fold-change e>2/<-2, the genes were considered to be differentially expressed.

**RRNA sequencing**

Fifty micron paraffin sections of tumor were used for RNA isolation using the RNaseasy FFPE kit per manufacturer’s instructions by Azenta Lifesciences (formerly Geneviz). Ribosome depletion was used to enrich for mRNA. The resultant RNA quality and quantity were determined by TapeStation and Qubit assays and used for library preparation. Libraries were sequenced on an Illumina HiSeq (150 base pair (bp) paired end reads) instrument (Azenta Lifesciences). Genome alignment was performed using STAR (V.2.7.0). Differential expression analyses were performed using DEseq2 (V.3.14) and computed immune cell densities were determined using CIBERSORT.

**RESULTS**

**Clinical data**

Relevant clinical data are summarized in table 1; safety results have been previously reported. All 12 patients with advanced unresectable melanoma had received anti-PD-1 therapy prior to lerapolturev while 6 of the 12
Table 1  Results of patient tumor flow cytometry analysis

| Pt no | Anti PD-1 resistance | Total no | Prior therapies/ prior anti- PD-1/prior anti CTLA-4 | CD3+Pre | CD3+Post | % change CD3+ | CD8+ Pre | CD8+ Post | % change CD8+ | CD4+ Pre | CD4+ Post | % change CD4+ | Current Status |
|-------|----------------------|----------|-----------------------------------------------------|---------|---------|-------------|---------|---------|-------------|---------|---------|-------------|----------------|
| 1     | Primary              | 1/yes/no | 11225                                               | 10996   | -2      | 3274        | 4419    | 35      | 2907        | 2414    | -17     |             | Anti-CTLA-4 plus anti-PD1, alive 3.2 years |
| 2     | Primary              | 3/yes/yes| 799                                                 | 903     | +13     | 324         | 389     | 17      | 152         | 176     | 16      |             | Died 23 months |
| 6     | Primary              | 3/yes/yes| 725                                                 | 211     |         | 199         |         |         |             |         |         |             | Died 11 months |
| 3     | Inadequate           | 2/yes/yes| 18923                                               |         |         | 5237        |         |         |             |         |         |             | Died 20 months |
| 4     | Secondary            | >10/yes/yes| 13932                                               | 9452    | -32     | 8771        | 4525    | -48     | 2304        | 1999    | -13     |             | Died 19 months |
| 5     | Secondary            | 4/yes/no | 8980                                                | 31697   | 252     | 1760        | 7968    | 352     | 3033        | 10228   | 237     |             | Anti-PD-1, progressed 25 months, alive 2.7 years |
| 7     | Secondary            | 2/yes/yes| 10557                                               | 55708   | 428     | 5054        | 24712   | 389     | 1642        | 9772    | 495     |             | Died 9 months |
| 10    | Secondary            | 1/yes/yes|                                                     |         |         |             |         |         |             |         |         |             | Died 13 months |
| 8     | Primary              | Yes/no   | 86395                                               | 148957  | 72      | 52799       | 89906   | 70      | 12387       | 28254   | 128     |             | Anti-PD-1, alive 2.5 years |
| 9     | Primary              | Yes/no   | 22376                                               | 14061   |         | 2681        |         |         |             |         |         |             | Anti-PD-1 alive 2.5 years |
| 11    | Secondary            | Yes/no   | 142502                                              | 13075   |         |             |         |         |             |         |         |             | Anti-PD-1 alive 2.2 years |
| 12    | Primary              | Yes/no   | 43591                                               | 98574   | 126     | 18382       | 56068   | 205     | 8299        | 36310   | 338     |             | Alive 2 years |

*These patients received anti-PD-1 within 30 days of PVRSIPO, therefore, confirmation of disease progression on anti-PD-1 prior to study enrolment at 4 weeks did not occur. Patients in gray are responders to lerapoltreuv.
patients had received both anti-PD-1 and anti-CTLA-4 therapy. Eleven of 12 patients also received anti-PD-1 therapy after lerapoltrev. Primary or secondary resistance was classified according to the Society for Immunotherapy of Cancer taskforce consensus. 

Briefly, primary resistance is best response of progressive disease (PD) or stable disease (SD) <6 months after ≥6 weeks of drug exposure while secondary resistance is best response of complete response, partial response, or SD >6 months after ≥6 months of drug exposure. Both definitions also include confirmatory scans at least 4 weeks after initial scan shows disease progression. After lerapoltrev treatment, patient 1 (primary resistance) received anti-PD-1 plus anti-CTLA-4 (no prior exposure to anti-CTLA-4) and continues without progression at 3.2 years. Two other patients with primary resistance died at 11 and 23 months after lerapoltrev. Of patients with secondary resistance (n=5), patient 11 had objective response to lerapoltrev and remains free of disease at 2.2 years, patient 5 restarted anti-PD-1 therapy after lerapoltrev and had disease progression after 25 months, while patients 4, 7, and 10 died after lerapoltrev. Finally, among patients (8, 9, and 12) with only prior anti-PD-1 therapy and who received anti-PD-1 therapy within 30 days before lerapoltrev (n=3), median progression-free survival (PFS) and overall survival (OS) after lerapoltrev is 2.3 years. Two of those patients restarted anti-PD-1 therapy (8, 9) and one received no additional treatment (12). As listed in table 1, all four patients with objective response to lerapoltrev (patients 8, 9, 11, and 12) have no evidence of disease and have current median PFS and OS of 2.3 years.

Flow cytometry and bulk RNA sequencing of tumors

Table 1 lists the absolute number of CD3+ and CD8+ cells in tumor specimens obtained before and after lerapoltrev as determined using flow cytometry. Of 17 samples included in our analysis with confirmed viable tumor on matched H&E, 200,000 events were analyzed in 12 samples whereas in 5 samples the number of cells shown in table 1 was normalized to 200,000 events. In table 1, the absolute number of cells is listed to account for differences in the total amount of viable tumor and tumor associated melanophages in the specimens.

Figure 1A,B shows CD4+ T cells and CD8+ T cell densities in tumors before and after lerapoltrev. Among seven patients with matched pre- and post-treatment tumors assessed via flow cytometry, two patients responded to lerapoltrev and five patients had no response to lerapoltrev. In three patients (non-responders), CD8+ T cells decreased whereas in four patients (two non-responders and two responders) CD8+ T cells increased, as listed in table 1 and shown in figure 1A. The mean increase in CD8+ T cells across 7 patients was 13,946 (p=0.0822, paired t-test). Across seven patients with matching lerapoltrev and postlerapoltrev samples, the mean increase in CD4+ T cells was 8998 (p=0.0522, paired t-test). The number of CD4+ T cells increased in two responders, increased in two non-responders, and appeared to be similar among the other three patients (figure 1B).

Next, we compared the differences in CD8+ and CD4+ T cells in the tumor at baseline (7–10 days prelerapoltrev) between three patients with median PFS (2.3 years) vs patients 7 with median PFS 1.6 months. In 3 patients who received anti-PD-1 within 30 days prior to lerapoltrev and have current median PFS 2.3 years, the mean number of CD8+ T cells in the pretreatment tumor was 28,414 (n=3) compared with a mean of 3519 in 7 patients with median PFS 1.6 months after lerapoltrev (p=0.0109), shown in figure 1C. Although the pretreatment tumor sample for patient 11, who has PFS 2.2 years was not evaluable, the post treatment tumor had 13,075 CD8+ T cells. Similarly, the mean pretreatment number of CD4+ T cells in patients with median PFS 2.3 years (n=3) was 7789 compared with 2269 (n=7) in patients with median PFS 1.6 months (p=0.0261), shown in figure 1C. These results were corroborated by immune cell enrichment as determined by CIBERSORT25 analysis of prerelapoltrev bulk RNA sequencing data of tumors (figure 1D, online supplemental figure S1). Patients 8, 9 and 12 (received anti-PD-1 within 30 days prior to lerapoltrev with PFS (2.3 years)) had significantly higher enrichment of CD8+ T cells (p=0.0134) and activated CD4+ memory T cells (p=0.0167), compared with seven patients with median PFS 1.6 months after lerapoltrev, shown in figure 1D. Patient 8 (PFS 2.5 years) had the highest number of CD8+ T cells as assessed by both flow cytometry and bulk RNA sequencing with patient 9 (PFS 2.5 years) having the second highest number of CD8+ T cell as assessed by both flow cytometry and RNA sequencing (online supplemental figure S1). Finally, bulk RNA sequencing revealed that patient 5 (PFS progressed at 25 months) had markedly higher levels of neutrophils in the TME compared with other patients (online supplemental figure S1).

DSP of tumors

To further examine the TME, we used NanoString DSP. Images after fluorescent staining (CD3, CD163, CD11c, DNA) and H&E stains for all prerelapoltrev specimens, are shown in figure 2. Each image also lists the number of CD3+ T cells in the TME as determined by flow cytometry. The H&E for patient 11 (PFS 2.2 months) showed necrotic tumor, and therefore, this sample was not included in all tumor analyses. This was thought to be a sampling error because the postlerapoltrev biopsy for patient 11 did have viable tumor. As shown in figure 2 and similar to flow cytometry and bulk RNA sequencing results, patients 8, 9, and 12 (received anti-PD-1 within 30 days prior to lerapoltrev with PFS 2.3 years) had markedly higher tumor infiltrating immune cells compared with patients 1–7 (median PFS 1.6 months).

Figure 3 details patient 2 and patient 8 as examples of DSP. Patient 2 (primary resistance, OS 23 months) had minimal immune infiltrate and the few CD3 cells present were in the periphery of the tumor, consistent with an
immune excluded profile. There were minimal differences between the prelerapolturlev and postlerapolturlev samples in patient 2 as assessed by DSP and flow cytometry (figure 3). Patient 8 (prior therapy was 6 cycles of anti-PD-1 therapy, with last dose of anti-PD-1 within 30 days of lerapolturlev, current OS 2.5 years) had more abundant immune cells compared with patient 2 (figure 3). There were no clear differences between the prelerapolturlev and postlerapolturlev biopsy in patient 8 (figure 3). For all patients with tumor available, slides were also fluorescently stained (DNA, CD3, CD11c, and CD163) and ROIs were selected for DSP analysis as shown in figure 3; ROIs were selected based on geometric location of tumor, examples of tumor regions are shown in figure 3F. Protein values below background were excluded and geometric regions intended to represent tumor that lacked MART-1 (melanoma marker) expression below threshold were also excluded. A heatmap of protein expression (13 proteins) in prelerapolturlev tissue relevant to the melanoma TME is shown in figure 3I. The values used for the heatmap are the mean SNR from at least two ROIs for each tumor sample. Online supplemental table 1 lists all SNR values used for comparison. Interestingly, patients 8, 9, and 12 (received anti-PD-1 within 30 days of lerapolturlev with median PFS 2.3 years) clustered together with higher but not significantly different mean HLA-DR expression (389.7 vs 21), higher mean CD45 expression (307.3 vs 36.4), higher mean CD8 expression (159 vs 9.6), and higher Ki-67 expression (99.6 vs 25.18) compared with prelerapolturlev tumor from patients with median PFS 1.6 months (n=7) figure 3I). This finding is also consistent with flow cytometry and bulk RNA sequencing where mean CD8+T cells from prelerapolturlev tumor in patients 8, 9, and 12 (received anti-PD-1 within 30 days prior to...
lerapolturev with PFS 2.3 years) were markedly higher compared with mean pretreatment CD8+T cells from pretreatment tumor in 7 patients with median PFS 1.6 months (28 414 vs 3519). Among 7 patients with matched prelerapolturev and postlerapolturev tumor available for DSP, there were no consistent patterns of change from prelerapolturev to postlerapolturev tumor in expression of the 13 proteins.

The limitation to selecting geometric tumor region is that some tumors have a visual pattern of infiltrative immune cells (patient 8; figure 3F and H), precluding geometric separation of immune cells vs tumor, while visually other tumors had regions with tumor cells only or had immune cells only in scattered regions (Patient 2; figure 3B and D). A comparison of the protein expression on immune cells only may provide insight into important differences about the properties of tumor infiltrating immune cells. Accordingly, we also performed protein selection on only CD3+ cells from tumors to determine the protein expression of CD3+ cells in the TME (figure 3). Online supplemental table 1 includes SNR for all proteins. Figure 3J shows expression of 12 proteins involved in T cell signaling in prelerapolturev tumor tissue from CD3+ cells only. It should be noted that expression of some markers such as MART-1 were present on CD3 cells (online supplemental table S1), suggesting there is some background protein expression not confined to CD3+cells. As expected, CD44 was high all for all patients since only CD3+ cells were selected for analysis. Patient 8, 9, and 12 (received anti-PD-1 therapy within 30 days of lerapolturev with median PFS 2.3 years) grouped together based primarily on high expression of CD8 compared with other patients. Among 7 patients with matched prelerapolturev and postlerapolturev tumor available for DSP, there were no consistent patterns of change from prelerapolturev to postlerapolturev tumor in expression of the 12 proteins on CD3+T cells.

We noted that while patients 8, 9, 12 (median PFS 2.3 years) appeared to have a more inflamed tumor profile primarily driven by CD8+T cells, other patients namely 5 (PFS 25 months) and 7 (death at 7 months) also had immune cell infiltrates within the TME visualized by fluorescent staining (figure 2). To explore differences in these immune infiltrates such as the presence of regulatory or other suppressor cells that may have led to the different clinical outcomes, we performed WTA on tumor from patients with immune infiltrates in tumor. FFPE tumor tissue sections from patients 8, 9, 12, (group 1) and patients 5 and 7 (group 2) were fluorescently stained for CD8 (teal), CD4 (red), CD11c (green), and DNA (blue, tumor) as shown in figure 4A, B. ROIs were then selected for areas enriched with CD8 and CD4 cells; WTA on cells only expressing the specific marker (either CD4 or CD8) was then performed as shown in figure 4A, B. At least two ROIs were selected for each cell type. Among 14442 human transcripts, we examined differences in
CD4+ cells and CD8+ cells between patients 8, 9, and 12 (group 1) and patients 5 and 7 (group 2). There were no significant differences in CD4+ transcripts between the two groups. Among CD8+ transcripts, five genes were significantly differentially expressed by more than twofold as shown in figure 4C, including higher HLA-DQA1, MIA (melanoma inhibitory activity), SERPINA3 (Serpin Peptidase Inhibitor, clade A member 3), KRTAP5-8 (keratin-associated protein 5–8), and XAGE1A (human cancer/testis antigens X-antigen family member 1) in group 1. These differences may reflect distinct activation status of CD8+ T cells from patients 8, 9, and 12 (median PFS 2.3 years) compared with CD8+ T cells from patients 5 (progressed at 25 months) and 7 (death at 7 months). Although this methodology analyzes only cells expressing the selected fluorescent marker (CD4+ or CD8+) there may also be background expression from tumor cells.

Analysis of peripheral blood

Immune subsets in blood were also analyzed at multiple time points. Patients with PFS 2.3 years after lerapolturev (patients 8, 9, 11, 12), which includes the three patients who had received anti-PD-1 therapy within 30 days before lerapolturev (8, 9, and 12) had significantly higher numbers of CD8+CCR7+CD45RA- T effector memory cells, but significantly lower CD8+PD-1+, and CD4+PD-1+ T cells compared with 8 patients with median PFS 1.6 months after lerapolturev (figure 5 and online supplemental figure S2). Patients 8, 9, and 12 (median PFS 2.3 years) had received a mean of 6 cycles of prior anti-PD-1 therapy prior to trial entry with last dose within 30 days before lerapolturev. No other correlations with response or changes prelerapolturev to postlerapolturev were noted.

In preclinical studies, PVSRIPO (lerapolturev) mediated antitumor efficacy by inducing type-I IFN dominant innate inflammation in myeloid cells via the cytoplasmic pattern recognition receptor MDA5.16,17 Moreover, recent evidence indicates that patient-specific differences in peripheral blood responsiveness to type I interferon associates with outcome after anti-PD-1 therapy in patients.20
We previously discovered that lerapoltrev infects and induces cytokine responses in monocytes, but not T/B cells in PBMCs after 24 hours of treatment. Thus, we next asked if peripheral capacity to induce antiviral inflammation after lerapoltrev infection associates with therapy outcome, possibly by reflecting the capacity of a patient’s innate immune system to respond to lerapoltrev treatment or by reflecting the status of peripheral innate immunity at the time of treatment. To this end, we developed a peripheral immune function assay that measured proinflammatory cytokine responses of pretreatment PBMCs after in vitro challenge with lerapoltrev or other innate stimuli to broadly probe innate immune status, including the TLR4 agonist LPS, the TLR1/2 agonist Pam3CSK4, and the TLR7/8 agonist R848 (figure 6; online supplemental figures S3 and S4). Since we observed higher levels of effector memory T cells in peripheral blood (figure 5) and tumor infiltrating T cells (figure 1) in patients living longer after lerapoltrev therapy, we also compared responses to T cell stimulation using CD3/CD28 ligation. The fold increase in cytokine production after 24 hours of stimulation with lerapoltrev over mock treatment is shown (figure 6; online supplemental figures S3 and S4). Patients with median PFS 2.3 years after lerapoltrev (patients 8, 9, 11, 12) including the 3 patients with known preraperoltrev inflamed TME who received anti-PD-1 therapy within 30 days before lerapoltrev (patients 8, 9 and 12), had significantly higher production of IFN-β, IFN-λ2, IFN-γ, and GMCSF compared with eight patients with median PFS 1.6 months after lerapoltrev. In addition, CXCL10 production was higher after LPS and anti-CD3/28 treatment in the four patients with PFS 2.3 years after lerapoltrev compared with eight patients with median PFS 1.6 months (online supplemental figures S3). No difference in baseline cytokine secretion (Mock treatment) was observed between patients with PFS 2.3 years vs PFS 1.6 months, indicating the aforementioned superior poststimulation cytokine levels in patients with PFS 2.3 years were due to differential sensitivity to in vitro stimulation (online supplemental figures S4). Together, peripheral preraperoltrev immune responsiveness (figure 6) with the increased T cell inflammation in preraperoltrev tumors and higher levels of effector memory CD8+T cells in preraperoltrev periphery of patients with PFS 2.3 years, imply that an active and/or proficient

Figure 4  NanoString Digital Spatial Profiling (DSP) of tumor tissue. NanoString DSP images after fluorescent staining on pretreatment tissue, CD8=teal, CD4=red, CD11c=yellow, and DNA=blue. ‘Low power’ = 2 mm field of view, 100 x total magnification. Rectangles indicate ROIs selected based on enrichment for cell type or tumor. (A) Low power image patient 12. (B) Low power image patient 7. (C) Volcano plot comparing ROI on CD8+ cells only between patients 8, 9, 12, 5 and 7. Differentially expressed genes appear in blue. ROI, regions of interest.
immune status at baseline in patients may result in favorable clinical outcomes.

**DISCUSSION**

The presence of tumor infiltrating lymphocytes (TILs) and an inflamed tumor in pre-checkpoint inhibitor therapy tumor (baseline) is associated with improved OS in melanoma and predictive of response to ICI therapy.\(^{30-32}\) Using three separate modalities—flow cytometry, DSP (protein and WTA), and RNA sequencing—we observed that three patients with advanced melanoma and a median PFS of 2.3 years had markedly elevated TILs in pre-lerapolturev tumor compared with seven patients with less abundant TILs in pretreatment tumors and median PFS 1.6 months. Although this observation aligns with previous findings that responses to immunotherapy preferentially occur in tumors with an ‘inflamed’ phenotype, our study is limited by a small sample size.\(^{31\text{a}}\) Furthermore, in matching pre-lerapolturev blood (baseline) from the same three patients with an inflamed TME, median PFS 2.3 years and a fourth patient with PFS 2.2 years (no pre-lerapolturev tumor available in the fourth patient), a more pronounced peripheral innate immune response to in vitro lerapolturev challenge was observed compared with eight patients with median PFS 1.6 months (including seven patients with non-inflamed pre-lerapolturev tumor). Findings in peripheral blood are also limited by small sample size. Among the eight patients with PFS 1.6 months, six patients have died while current PFS and OS among the other four patients is 2.3 years.

Determining clinical and radiographic responses to anti-PD-1 therapy can be complicated by delayed responses and pseudoprogression.\(^{36\text{a}}\) Guidelines that more precisely define anti-PD-1 failure predated our

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**Figure 5**

Flow cytometry analysis of immune subsets in peripheral blood. PBMCs were analyzed for T cell memory populations and PD1 expression on CD8+ and CD4+ conventional T cells as shown at the pretreatment time point. Patients in red have median PFS of 2.3 years after lerapolturev and patients in black have median PFS 1.6 months after lerapolturev. Data brackets indicate mean ±SEM; p values are from unpaired t-test. PBMCs, peripheral blood mononuclear cells; PFS, progression-free survival.
study design and as such five patients received anti-PD-1 therapy within 30 days of lerapopture, although clinically these patients appeared to be progressing. Three of those patients (8, 9, and 12) had the highest number of CD8+T cells (mean 28,414) in prelerapopture tumor (tumor biopsied within 30 days of anti-PD-1 therapy prelerapopture) compared with mean value of 3509 (n=7) in other patients. This was consistent with bulk RNA sequencing and DSP analysis. In addition, patient 11 with PFS 2.2 years, had a higher number (13,075 CD8+T cells) in postlerapopture tumor. Together, those four patients (8, 9, 11, 12) continue to have prolonged PFS (median 2.3 years vs 1.6 months median PFS for the other seven patients). Interestingly, patient 9’s (PFS 2.5 years) prelerapopture non-treated lesion also had marked tumor inflammation while patient 1’s (SD after initiation of anti-CTLA-4 plus anti-PD-1) prelerapopture non-treated lesion had little immune infiltrate (figure 3J). Therefore, it is possible that anti-PD-1 therapy contributed to disease control for patient 8, 9, and 12, (median PFS 2.3 years) since the last dose of anti-PD-1 therapy was within 30 days and the TME of those patients had a more inflamed tumor profile prior to lerapopture. However, lerapopture may have provided additional support of inflammation and remodeling of the TME. Recent data suggest that there are T cell infiltrated tumors that are resistant to anti-PD-1 therapy, possibly due to lack of coexisting proinflammatory cytokines. Oncolytic viruses like lerapopture may provide this type of inflammation which can lead to a clinical benefit as we observed in four patients on the study.

Use of spatial biology in our study in addition to flow cytometry and bulk RNA sequencing highlighted interesting aspects about studying the TME. Intratumoral heterogeneity was observed which is most apparent when viewing DSP images, which preserves spatial biology. Some patients had portions of tumor with abundant TILs, some had TILs only on the tumor periphery, and some patients had TILs arranged in an infiltrative pattern. Those relationships are not as easily captured using flow cytometry or bulk RNA sequencing. However, flow cytometry can provide more detailed information about live cells, and bulk RNA sequencing evaluates a larger spectrum of cell types. Although CD8+ T cell infiltrated tumors are a known predictor of response to anti-PD-1 therapy, about 30% of patients with CD8+ T cells in the tumor will still not respond to checkpoint therapy. WTA of CD8+ T cells among two groups of patients, both with CD8+ T cells present in tumor but with divergent outcomes revealed differences in the CD8+ T cells that may in part explain the divergence. The higher HLA-DQA1 on CD8+ T cells from patients with prolonged PFS suggests robust T cell activation. Spi2A, is encoded by SERPINA3. Spi2A enhances the initial development of memory CD8+ T cells and SERPINA3 was markedly increased on CD8+ T cells from patients with prolonged PFS.

Given the intratumoral heterogeneity and limitations of biopsies which do not evaluate the entire tumor, we also conducted blood analysis. Identification of a blood-based biomarker could also be a useful approach to evaluate therapy efficacy in larger clinical studies given that blood is easier to obtain, non-invasive, and facilitates longitudinal analysis of changes in the immune system. Previous reports indicate that anti-PD-1 therapy induces proliferation of CD8+ PD-1+ and CD4+ PD-1+ T cells in peripheral blood with the increase peaking at 1 week after initiating anti-PD-1 and then decreasing thereafter despite continuous anti-PD-1 treatment. Therefore, the finding that three of four patients with prolonged PFS (2.3 years) in our study had lower CD8+ PD-1+ and CD4+ PD-1+ T cells is not surprising since three of four patients with prolonged...
PFS had received an average of six cycles of anti-PD-1 with last dose within 30 days prior to lerapolturev while only two of eight patients with poor PFS had received anti-PD-1 within 30 days.

Since anti-PD-1 therapy works by expanding functional effector T cells in the TME and lerapolturev also recruits peripheral blood innate immune cells in the TME, we investigated if immune function in blood could predict presence of immune cells in the TME or ability to recruit peripheral cells into the tumor. Three patients who received anti-PD-1 within 30 days of lerapolturev, who had an inflamed prerapolturev TME as noted by flow cytometry, DSP, and RNA sequencing, and had PFS 2.3 years, appeared to elicit a robust response in peripheral blood as measured by cytokine production after in vitro stimulation with lerapolturev (and in some cases LPS and anti-CD3/CD28 stimulation) compared with eight patients with median PFS 1.6 months. A fourth patient (11) with no prerapolturev tumor available (but who had an inflamed post lerapolturev TME), with PFS 2.2 years also mounted a robust response in peripheral blood after in vitro stimulation with lerapolturev. Of note, all four patients with prolonged PFS (2.3 years) and robust innate immune response had only received anti-PD-1 therapy prior to lerapolturev while all other patients were more heavily pretreated (table 1). This type of analysis may be useful in assessing peripheral innate immune responsiveness/competence and may predict which therapies can effectively recruit immune cells into the TME for an individual patient. Four patients with the most robust cytokine response had the highest levels of T cells in the TME.

A current trial, LUMINOS-102 (lerapolturev with or without immune checkpoint blockade) in patients with advanced PD-1 refractory melanoma (NCT04577807) is ongoing. Patients will have confirmed progression on anti-PD-1 therapy before enrollment. In treatment naive patients, final results of MASTER-KEY-265 suggested that the intratumoral oncolytic virus, talimogene laherparepvec plus pembrolizumab did not significantly improve PFS or OS compared with placebo plus pembrolizumab in a large phase III randomized study. Therefore, intratumoral therapies may be more relevant for selected patients not initially responding to checkpoint therapy as is being explored in LUMINOS-102. In addition, the mechanisms of lerapolturev are unique and efficacy may be different.

In conclusion, patients with a prerapolturev inflamed TME and increased sensitivity to in vitro lerapolturev challenge in prerapolturev peripheral blood had prolonged median PFS (2.3 years) after treatment with lerapolturev compared with patients with median PFS of 1.6 months, who had prerapolturev ‘cold’ tumors and lower cytokine response in prerapolturev peripheral blood. All four patients with prolonged PFS had also received anti-PD-1 therapy (three of 4 within 30 days of lerapolturev) and no additional systemic therapies. Given this observation in our pilot study in 12 patients, future studies will test if immune response measured in the periphery (blood) may be important in predicting ability to recruit immune cells into the TME in a larger cohort of patients.

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