Pre-treatment tumor neo-antigen responses in draining lymph nodes are infrequent but predict checkpoint blockade therapy outcome

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ABSTRACT

Immune checkpoint blockade (ICPB) is a powerful effective cancer therapy in some patients. Tumor neo-antigens are likely main targets for attack but it is not clear which and how many tumor mutations in individual cancers are actually antigenic, with or without ICPB therapy and their role as neo-antigen vaccines or as predictors of ICPB responses. To examine this, we interrogated the immune response to tumor neo-antigens in a murine model in which the tumor is induced by a natural human carcinogen (i.e. asbestos) and mimics its human counterpart (i.e. mesothelioma). We identified and screened 33 candidate neo-antigens, and found T cell responses against one candidate in tumor-bearing animals, mutant UQCRNC. Interestingly, we found a high degree of inter-animal variation in the magnitude of neo-antigen responses in otherwise identical mice. ICPB therapy with Cytotoxic T-lymphocyte-associated protein (CTLA-4) and α-glucocorticoid-induced TNFR family related gene (GITR) in doses that induced tumor regression, increased the magnitude of responses and unmasked functional T cell responses against another neo-antigen, UNC45a. Importantly, the magnitude of the pre-treatment draining lymph node (dLN) response to UNC45a closely corresponded to ICPB therapy outcomes. Surprisingly however, boosting pre-treatment UNC45a-specific T cell numbers did not improve response rates to ICPB. These observations suggest a novel biomarker approach to the clinical prediction of ICPB response and have important implications for the development of neo-antigen vaccines.

Introduction

Immunotherapies such as anti-cytotoxic T lymphocyte associated antigen (CTLA)-4 and anti-programmed death (PD)-1 antibodies can cause long term, durable tumor regression in some cancers. Tumor mutation burden correlates with immune checkpoint blockade (ICPB) outcomes in multiple studies\(^1\)\(^-\)\(^3\) and as such, favorable responses to ICPB have been reported in patients with tumors caused by carcinogen exposure\(^1\) with consequently high-predicted numbers of neo-antigens.\(^4\)\(^-\)\(^3\) However, the actual role of neo-antigens in these responses remains uncertain.

Neo-antigens, altered peptides derived from somatically mutated tumor DNA, can be predicted from whole exome and RNA sequencing in combination with \textit{in silico} MHC binding affinity algorithms. However, \textit{ex vivo} testing often finds that only a small proportion of predicted neo-antigen candidates are immunogenic.\(^4\)\(^-\)\(^7\) Whilst this can be partly attributed to limitations in current prediction methods and neo-antigen presentation, the absence of neo-antigen responses could be due to the failure of the immune system to generate a detectable response in tumor bearing individuals. A recent study demonstrated that tumor neo-antigen specific T cells can be expanded from HLA matched healthy donor lymphocytes, but not from cancer patient lymphocytes.\(^7\) This suggests that some tumor bearing individuals have a limited capacity to generate a detectable neo-antigen response which may be caused by a negative regulatory tumor microenvironment.

ICPB removes some of the negative regulatory pressures exerted on T cells. It has been observed that ICPB increases the magnitude of T cell responses against tumor neo-antigens, enabling detection of neo-antigen specific immune responses not detectable prior to treatment.\(^1\)\(^,\)\(^8\)\(^,\)\(^9\) To explore this notion and assess whether ICPB response rates could be improved we used a mouse mesothelioma model, AB1-HA.

This model was chosen for these studies because it is one of the few tumor models that is induced by the relevant human carcinogen (i.e. asbestos) and exhibits histological, clinical, immunological and mutational characteristics similar to the equivalent human cancer, mesothelioma.\(^10\) In addition, AB1-HA is susceptible to immunotherapy\(^11\)\(^-\)\(^14\) despite not having a high mutation burden,\(^15\)\(^,\)\(^16\) as preliminary studies are now suggesting for the human counterpart.\(^12\)\(^,\)\(^17\) AB1-HA has two known, tractable tumor antigens, a previously described neo-antigen, UQCRNC\(^2\)\(^8\) and hemagglutinin\(^2\)\(^9\) previously transfected into the cell line as a model neo-antigen.\(^2\)\(^9\)
We hypothesized that ICPB would increase the magnitude of these neo-antigen specific T cell responses as well as unmasking responses to additional neo-antigens from in silico predicted candidates. Furthermore, we examined whether pre-existing immune recognition of neo-antigens reflected response rates to ICPB. This has been hard to determine directly because of the inherent variability in response to ICPB which of course can only be assessed after therapy. In order to overcome this limitation, we used a recently developed dual-tumor model that enables the status of the tumor to be assessed prior to therapy and this crucial question to be addressed.

Materials and methods

Mice

Eight to 10-week old female BALB/c and C57J/BL6 mice were purchased from the Animal Resource Center, Murdoch, Australia and maintained under standard specific pathogen-free housing conditions at the Harry Perkins Institute of Medical Research. All animal experiments were conducted with the approval of Harry Perkins Institute of Medical Research Animal Ethics Committee.

Murine mesothelioma and lung cancer models

The murine mesothelioma cell line AB1 was previously generated as described.\(^{20}\) H1N1/PR8 influenza hemagglutinin HA was transfected as a model neo-antigen to generate AB1-HA.\(^ {19}\) Cells were maintained as previously described.\(^ {18}\) The AE17 murine mesothelioma cell line was established in 2003 by exposing C57BL/6j mice to crocidolite asbestos.\(^ {21}\) The Line1 murine alveolar carcinoma (Line1) was established in 1974.\(^ {22}\) Original stocks were obtained from Professor Najat Eglêmeze, University of Louisville (KY, USA). Line1 was cultured with DMEM (Gibco) containing 20mM HEPES and supplemented with 10% FCS. Cells were used at below 20 passages for experiments and were confirmed to be negative for Mycoplasma spp by PCR.

5 x 10\(^7\) tumor cells were injected subcutaneously in the flank of mice (right flank for single tumor model, bilateral flanks for dual-tumor model). Mice were euthanized at indicated timepoints. Ipsilateral axillary and inguinal tumor draining lymph nodes (DLN) were studied because they are the primary location of neo-antigen cross-priming in this and other murine models,\(^ {18}\) optimizing our chances of detecting significant and biologically relevant differences.

Combination immune checkpoint blockade

Mice bearing AB1/AB1-HA were administered intraperitoneally (ip) with 100μg anti-CTLA-4 (clone 9H10; Bioceross; Netherlands) and 50μg anti-GITR (clone DTA-1; BioXCell, NH, USA) antibodies in a total volume of 200μL PBS at indicated time points. Animals were dosed at concentrations previously shown to induce AB1-HA regression\(^ {12}\) when tumor are between 16 mm\(^2\) and 20 mm\(^2\) in size. An equivalent volume of PBS was injected ip in control animals. Mice were randomized to treatment or control arms and tumor measurements were made blinded to treatment group.

\(\text{Tumor and DLN resection}\)

We utilized a version of the dual-tumor model developed by Lesterhuis et al.,\(^ {13}\) in which bilateral tumors progress or regress synchronously and symmetrically during therapy with ICPB. For experiments with bilateral tumors, the right flank tumor and its corresponding inguinal lymph nodes were surgically resected at day 11 as previously described.\(^ {23}\) The following day, mice received 100μg anti-CTLA-4 and 50μg anti-GITR ip and growth of the left flank tumor was monitored.

\(\text{Mutation calling and neo-antigen prediction}\)

AB1/AB1-HA whole exome sequencing (WES) and RNA sequencing (RNA-seq) data were reanalyzed for this study.\(^ {18}\) WES and RNA-seq of Line1 and AE17 were performed under contract by Australian Genome Research Facility. DNA and RNA were extracted from in vitro cultured cells. DNA was processed with SureSelectXT Mouse All Exon kit (Agilent, Australia) and RNA with Illumina’s TrueSeq Stranded mRNA library preparation protocol. Paired-end sequencing (2x100bp) was performed on the Illumina HiSeq 2500 where >50 million total reads per sample were generated.

WES data were aligned to the updated murine reference genome mm10 (UCSC mm10/NBNI GRCm38) using Burrows-Wheeler Aligner.\(^ {24}\) RNA-seq data were aligned to mm10 using Hisat2 v2.04.\(^ {25}\) Expression analysis of aligned RNA-seq data was performed with Stringtie v1.3.\(^ {26}\)

Somatic single nucleotide variants (SNVs) were identified from WES using both Varscan 2.3.6\(^ {27}\) and Somatic sniper 1.0.5.\(^ {28}\) High-confidence SNVs as defined by Varscan (at least 10% mutant reads in tumor, < 5% in normal and a one-tailed Fisher’s exact test P-value < 0.07) and Somatic sniper (variant supporting reads with an average mapping quality of ≥ 40 and a somatic score of ≥ 40) were pooled. Variants were annotated with Annovar\(^ {29}\) and SNVs were filtered to identify non-synonymous, exonic variants.

The MHC-I binding affinity of mutated sequences were predicted as previously described\(^ {18}\) to respective MHC-1 haplotype. MHC-I binding affinity of AB1/AB1-HA and Line1 SNVs were predicted for H2-Dd, H2-Kd and H2-Ld, and AE17 SNVs for H2-Dd and H2-Kb. Briefly, predicted MHC I binders were selected based on their relative ranking in NetMHCpan 2.8 where the top 0.5% of ranked peptides were considered strong binders, top 2% ranked peptides weak binders, and those ranked > 2% non-binders.\(^ {30}\) The expression of predicted strong and weak binders was confirmed with RNA-seq data and candidates with > 1 FPKM and the mutant allele expressed were selected.

A number of potential MHC binders can be predicted from a peptide sequence. The epitope with the top predicted binding affinity for each variant sequence was selected for peptide synthesis. Short peptides of 8 to 11 amino acid length were synthesized. Long peptides (27 amino acids long with the point mutation at position 13) for 9 of the top 10 expressed candidates were synthesized. Peptides (Mimotopes, VIC, Australia) were dissolved in 1% DMSO (Sigma Aldrich, NSW, Australia) and Ultrapure™ DNase/RNase-free distilled water (Invitrogen, VIC, Australia) to a concentration of 1 mg/mL and stored at −20°C.
**IFN-γ ELISPOT**

IFN-γ production in response to neo-antigen was examined by ELISPOT. Single-cell suspensions of the dLN were prepared by mechanical disruption. Tumors were cut into 1 mm² pieces with a scalpel, and digested with a mouse tumor dissociation kit (Miltenyi Biotec, NSW, Australia) on the GentleMACSTM Octo Dissociator (Miltenyi Biotec). Tumor cell suspensions underwent CD8+ T cell selection with CD8 Microbeads (Miltenyi Biotec) as per manufacturer’s instructions. 1 × 10⁵ freshly isolated dLN cells or 1 × 10⁴ post-CD8-selection tumor infiltrating lymphocytes (TILs) were incubated per well in a microtiter plate and stimulated with 1 μg/mL of individual peptides. Peptide pools consisting of 8–25 peptides, with each individual peptide at a final concentration of 1 μg/mL, were also tested. As a positive control, 0.5 μL/well of Dynabeads® Mouse T-activator CD3/CD28 beads (Thermo Fisher Scientific, NSW, Australia) were used. The volume of beads used was empirically calculated to enable enumeration of the positive control. All ELISPOT assays were performed in duplicate and results presented as IFN-γ Spot Forming Units (SFU) per 10⁵ cells, unless stated otherwise. IFN-γ SFU made in response to mutant peptides were compared to those made in response to corresponding wild type peptides. In situations where the wild type peptide was unavailable or not applicable (such as for HA responses), the media only negative control was used as a comparison.

**In vivo cytotoxic t lymphocyte (CTL) assay**

Splenocytes were prepared from non-tumor bearing BALB/c mice and resuspended at 1 × 10⁷ cells per mL in RPMI containing 2% fetal calf serum. Target splenocytes were pulsed with either 1 μg/mL HA518-526 (IYSTVASSL), UNC45a730-738 mutant (IYEVRRLSV), UQCRCC2405-413 wildtype (SYMPPTSTVL) or UQCRCC2405-413 mutant (SYMAPPTSTVL) peptides for 1 h at 37°C. After washing, cells pulsed with two different peptides were labeled with 2.5 μM of either CFSE (Invitrogen, VIC, Australia) (CFSE®) or Violet-Tag (Biolegend, CA, USA) (Violet®) for 10 min at 37°C. Control unpulsed cells were labeled with 0.25 μM CFSE (CFSE®) or Violet-Tag (Violet®). Cells were resuspended to 1×10⁶/mL. 1 × 10⁷ peptide pulsed cells and 1 × 10⁷ unpulsed cells were injected intravenously into tumor-bearing mice treated with ICPB. dLNs were collected 18–20 h later and single-cell suspensions were prepared. Cells were stained with anti-CD8a (clone 53.67, Biolegend, CA, USA) for 20 min at room temperature. Cells were washed and analysis was performed on a BD LSRSFortessa (San Jose, CA, USA). Data were analyzed using FlowJo software (Tree Star, Ashland, OR, USA). Percent-specific lysis was calculated as: (1−(R_{BALB}/R_{ABIH-BALB}) × 100, where R = %CFSE°/%CFSE°.

**Neo45a mutant immunization**

UNC45a mutant synthetic long peptide (MTFPGERIYEYVRSLVSLHLSCSGLQ) was emulsified in Montanide™ ISA 51 VG (Seppic, Paris, France). A total of 40 μg peptide per mouse was made up to 100 μL volume with PBS and emulsified in a 1:1 ratio with 100 μL Montanide in accordance to the manufacturer’s instructions. Animals were injected with 100 μL of the vaccine subcutaneously at the base of the tail on both the left and the right flank (equivalent to 20 μg peptide per flank).

**Statistical analysis**

Data were analyzed with Graph Pad Prism Software (GraphPad software, CA, USA) with results presented as mean ± SD. Paired student’s T-test was used to compare IFN-γ responses of individual mice to mutant peptide or controls. A p value of <0.05 was considered significant and indicates a positive response to the test peptide. To assess the variability of responses between mice, results were presented as median ± IQR and the Levene’s test was used to determine the homogeneity of the neo-antigen responses. One-way ANOVA with Dunnett’s multiple comparison was performed to compare long-peptide responses with media negative control. Linear regression was performed to determine correlations. Fisher’s exact test was used to compare UNC45a response in treated and untreated mice. The effect of immunotherapy on the magnitude of neo-antigen T-cell response was analyzed in a mixed model ANOVA. The background effect was adjusted in the analysis as a covariate, the treatment was entered as a fixed effect and the experiment was entered as a random effect. The dependent variable was computed as the difference between mutant and wildtype peptide response. The log rank test (Mantel Cox) was performed to compared survival between different treatment groups.

**Results**

**Few in silico predicted neo-antigen elicit an immune response**

In AB1, a total of 2,697 high-confidence SNVs were indentified while there were 1,047 SNVs in AE17 and 3,620 in Line1. Of these, 434, 14 and 361 non-synonymous exonic SNVs were predicted to be MHC class I binders where 78, 13 and 104 were expressed in the three cell lines, respectively.

In our previous study, we described immune responses against 1 out of 20 tested candidate neo-antigens in the murine mesothelioma, AB1. In the present study, we increased the number of tested candidate neo-antigens. A total of 16 new candidates were tested, 8 that were identified previously but not tested and 8 new candidate neo-antigens that were identified. Of these 16 new peptides, 6 were predicted to be strong MHC-I binders, and the other 10 were predicted to be weak MHC-I binders (Supplementary table I).

We screened for ex vivo IFN-γ production in response to a total of 33 neo-antigen short peptides (17 previously tested and 16 new candidates) from AB1 murine mesothelioma, 31 top candidates based on predicted MHC class I binding affinity from Line1 lung carcinoma and 13 from AE17 murine mesothelioma. There was no significant response against any of the 33 mutant peptides in the dLN of AB1-HA bearing animals (Figure 1a and b) yet strong, spontaneous responses to mutated UQCRCC2 and HA peptides were again detected (Figure 1b). No responses were detected to the 31 neo-antigen candidates in Line1 lung carcinoma (Figure 1c) and 13 neo-antigen candidates in AE17 murine mesothelioma (Figure 1d). Tested peptide sequences are listed in supplementary...
figure II. Thus, a measurable immune response was only detected in AB1-HA tumor bearing animals to one of the panel of in silico predicted neo-antigens.

Variability in immune responses to neo-antigens

Neo-antigen responses in outbred humans are variable. To examine the stochasticity of neo-antigen responses we studied inbred mice that are genetically identical, handled identically and bear identical tumors, examining the range of responses to these neo-antigens. Substantial variability in immune responses against HA and mutant UQCRC2 peptides were observed between AB1-HA bearing animals (median response to HA: 17 SFU/10^5 cells, IQR 7–52; Mutant UQCRC2: 22 SFU/10^5 cells, IQR 11–43) (Figure 2a). Variability in IFNγ SFU was also observed in unstimulated or wild-type UQCRC2 stimulated wells, and mice with high levels of background reactivity were generally more likely to respond to neo-antigen peptide. The number of IFNγ SFU in response to HA and UQCRC2 mutant peptide correlated with each other (r^2 = 0.7308, p < 0.0001) (Figure 2b). There was no association between increasing tumor size and the magnitude of the neo-antigen responses (HA r^2 = 0.03294, p = 0.1192; Mutant UQCRC2 r^2 = 0.01283, p = 0.3365). Thus neo-antigen responses to identical tumors varied hugely between identical animals with some mice generating strong neo-antigen specific T cell responses but others generating none at all.

One new neo-antigen response was detected following ICPB

As there were minimal detectable immune responses against our panel of candidate neo-antigens, we hypothesised that ICPB would increase the magnitude and substantially broaden the range of neo-antigen responses, as suggested by some studies, albeit in outbred humans. AB1-HA bearing animals were treated with a combination of anti-CTLA-4 and anti-GITR, which results in reliable immune-mediated tumor regression. We screened for responses in the dLN against neo-antigen candidate peptides of various lengths in treated animals (Figure 3a).

We first tested for dLN responses against short peptides of the 33 neo-antigen panel and found significant responses against HA and mutant UQCRC2 peptides in both treated and untreated mice. No responses were observed to any of the other short peptides (Figures 3b and c).

CD8+ T cells typically respond to 8-11mer short peptide epitopes in an ELISPOT assay, whereas both CD4+ and CD8+ T cells can respond to 27mer long peptides in an ELISPOT assay with antigen-presenting cells. To further explore responses to our neo-antigen candidates, we tested long peptides encoding the top nine expressed neo-antigen candidates (listed in Supplementary table III). Responses to mutant UQCRC2 long peptide were again detected and seen in both untreated and treated mice (Figure 3d, e). We detected
immune responses to an additional neo-antigen, mutant UNC-45 homolog A (UNC45a) in the dLN of treated animals, but not untreated animals (Figures 3e and g). All other neo-antigen long peptides failed to elicit any IFNγ production regardless of treatment. Thus ICPB unmasked one additional neo-antigen rather than many as we had hypothesized.

Minimal UNC45a epitope is not the strongest predicted binder

We did not detect responses to the 8mer mutated UNC45a peptide (IYEVVRSL), which was predicted to be the strongest MHC H2-Kd binder with a binding affinity of 194 nM, in our peptide pool (Figure 3b). We tested two more predicted minimal epitopes within the mutant UNC45a sequence, and observed reactivity to the 9mer IYEVVRSLV (H2-Kd, 364nM), but not RSLVSLHL (H2-Dd, 7139nM)(Figure 3g). This suggests that IYEVVRSLV is a minimal mutant UNC45a epitope. Thus selection based on predicted binding affinity alone could be insufficient in selecting an immunogenic neo-antigen epitope.

In vivo cytotoxic activity of neo-antigen specific T cells in DLN

We examined the cytotoxic function of neo-antigen specific T cells within the dLN in ICPB treated mice and found significantly higher HA-specific CTL activity in the dLN (40 ± 15%) compared to the non-dLN (12 ± 15%) (p = 0.0018; n = 5), strongly suggesting that, as predicted, the dLN was the dominant site of neo-antigen specific CTL activity (Figures 3h and 3i). A significantly higher UQCR2 specific CTL activity was also observed in the dLN (36 ± 18% vs 20 ± 12%; p = 0.0107). The UNC45a specific CTL activity was not significantly different between the two sites (22 ± 20% vs 6 ± 8%; p = 0.057). This supports previous observations that the examination of the dLN cells can provide a useful profile of the neo-antigen reactivities.18,33

Icpb-induced neo-antigen specific T cells are not excluded from the tumor

We have previously shown that not all cross-primed neo-antigen specific CD8+ T cells in the dLN enter the tumor microenvironment,34 raising the concern that the typical studies of neo-antigen specific T cells obtained from tumor tissue rather than dLN may miss some of the relevant responses. When we investigated this using the neo-antigens described above, we found that ICPB immunotherapy increased the magnitude of all three neo-antigen T cell responses in the dLN and in the tumors. Responses to HA, mutant UQCR2 and UNC45a in the dLN of treated mice were significantly higher compared to untreated mice (Figure 4a-c) (HA, 96 ± 101 vs 10 ± 35 SFU/10^5, p = 0.03; UQCR2, 73 ± 71 vs 19 ± 22 SFU/10^5 p = 0.021; UNC45a, 83 ± 88 vs 15 ± 28 SFU/10^5, p = 0.03) and this increase in magnitude was
also observed in the tumor (Figure 4d-f) (HA, 107 ± 68 vs 34 ± 32 SFU/10^4 CD8^+ TILs, p = 0.002; UQCRC2, 172 ± 97 vs 53 ± 44 SFU/10^4 CD8^+ TILs, p = 0.014; UNC45a, 159 ± 86 vs 18 ± 18 SFU/10^4 CD8^+ TILs, p < 0.001). ICPB increased the proportion of individual mice that made a significant response to UNC45a in both the dLN (15% to 73%) and tumor (20% to 90%) (Table 1). A positive response was defined as a significant difference (p < 0.05) between indicated neo-antigen and media control (for HA) and wild-type peptide (for UQCRC2 and UNC45A) using paired student’s T-test.

Two-tailed Fisher’s exact test was performed to compare between treated and untreated mice.

**Strong dLN responses to UNC45a prior to ICPB therapy predict successful treatment outcome**

As the endogenous neo-antigen response varied between identical animals, we hypothesized that those with stronger mutant UQCRC2 and UNC45A ELISPOT responses would successfully respond to ICPB therapy. To determine if pre-treatment neo-antigen responses correlated with therapy outcome, we utilized an established bilateral tumor model. The symmetry in this model allowed us to resect a tumor and dLN before treatment to determine the level of neo-antigen response, whilst leaving the other tumor as a readout of subsequent therapy response (Figure 5a and b). Around half of the mice treated with ICPB responded to treatment as evidenced by the complete regression of tumor in 12 out of 22 mice. Pre-treatment dLN UQCRC2 response was significantly higher in responders (n = 12, 17 ± 12 SFU/10^5) compared to non-responders (n = 10, 7 ± 5 SFU/10^5, p = 0.0182) (Figure 5c). Similarly, the UNC45a response was significantly higher in the responders (24 ± 14 SFU/10^5 vs 5 ± 2 SFU/10^5, p = 0.0004) (Figure 5d). This suggests that the variability of neo-antigen immune responses between mice is biologically significant and responses to some neo-antigens could predict subsequent response to ICPB therapy.

**Boosting unc45a immune responses did not improve ICBP outcomes**

As low UNC45a specific T cell responses pre-treatment was closely associated with failure to respond to ICPB therapy, we hypothesized that ICBP outcomes would be improved by increasing the magnitude of pre-treatment UNC45a responses through peptide immunization (Figure 6a). Administration of UNC45a synthetic long peptide in montanide significantly increased the immune response to UNC45a in vaccinated mice (n = 19, 34 ± 21 SFU/10^5) compared to both unvaccinated mice (n = 15, 8 ± 8 SFU/10^5, p < 0.0001) and to mice that received montanide alone (n = 18, 13 ± 15 SFU/10^5, p = 0.0009) (Figure 6b). There was no...
difference in the response to UQCRC2 neo-antigen which was not included in the vaccine (Figure 6b).

Importantly, UNC45a vaccination alone did not have any effect on tumor growth (Figure 6c). Of mice that received both montanide and ICPB, 20% (2/10) had a complete response (Figure 6d). Mice that received ICPB alone had 20% (2/10) partial responders where tumor growth was hampered after treatment but tumors eventually grew out (Figure 6e). Of those that received both the UNC45a vaccine and ICPB, 30% (3/10) were complete responders (Figure 6f). As before, ICPB treatment significantly improved survival \( p = 0.0101 \) compared to untreated mice (Figure 6g). Whilst there were some long-term survivors in mice that received combination vaccine and ICPB therapy, the effect size was small and there was no significant benefit in survival compared to those that received ICPB alone \( p = 0.2188 \). Again, mice that responded to ICPB had a higher UNC45a response pre-treatment compared to non-responders \( 37 \pm 20 \text{ SFU}/10^5 \) vs \( 11 \pm 12 \text{ SFU}/10^5 \), \( p = 0.0016 \) (Figure 6g). This result suggests that a strong UNC45a response could predict a therapeutic response to ICPB treatment but it is not a sufficient individual component of an effective ICPB-induced response.

**Discussion**

Tumor mutations can be targets of anti-tumor immunity, and mutational load has been associated with response to ICPB in the clinic. Whilst few spontaneous immune responses against genomically predicted neo-antigens have been detected, the responses appear to be important in immune-mediated rejection of tumor. Thus these neo-antigen responses could represent predictors of response to therapy.
In this and other studies, few sequencing derived neo-antigen candidates elicit immune responses in the presence of tumor. Most published murine neo-antigen reactivities have been identified postvaccination.39, 40 Our previously identified UQCRC2 neo-antigen in AB1-HA18 and three neo-antigens identified in a murine colon adenocarcinoma model38 were detected in tumor-bearing mice without treatment. The paucity of neo-antigens identified in this setting could be partly attributed to the limitations of current epitope prediction algorithms and selection, but also to tumor-associated immune suppression. Relaxing the immune suppressive environment in our model by ICPB increased the magnitude of immune responses against the endogenously recognized neo-antigen UQCRC2 by 3.5-fold and enabled immune responses to a previously undescribed neo-antigen, mutant UNC45a, to be detected. Somewhat surprisingly, there were minimal responses against the rest of the candidate neo-antigens. This suggests that ICPB increased the dynamic range of immune responses which improved our ability to detect the neo-antigen response, a finding that has been observed in other studies.8, 9, 35 This result also supports the hypothesis that immune responses to neo-antigens in tumor-bearing hosts are attenuated and yet partially reversible by CTLA-4 treatment.41

In this study, we observed significant inter-mouse variation in neo-antigen reactivities before ICPB therapy. Such variability has previously been reported in in-bred mice bearing MC-38 murine colon adenocarcinoma.38 There are many potential factors underlying such T cell response heterogeneity, including stochastic variability in the TCR repertoire in otherwise genetically identical mice. We considered it unlikely that the microbiome was a cause for variability as it has been shown that animals housed in the same cage and within the same animal facility have little variation in microbiomes.42 The issue of variable immune responses is much harder to study in outbred humans. Recent melanoma studies have shown that individual patients have variable responsiveness to neo-antigen in terms of both the number of responses and their magnitude before and after personalized neo-antigen vaccination.5, 6 It is impossible to tell if this variability is related to the host immune responsiveness or other factors to do with the candidate selection, vaccine or T-cell repertoire. Our data suggest that an inherent variability in neo-antigen responses may underlie at least some of this interpatient variability. Understanding the determinants that underlie this response variability may enable the selection of a therapeutic strategy that is useful for generating an anti-tumor response in individual patients.

One of the major issues confronting cancer immunologists at the moment is to determine which patients will respond to ICPB. Tumor mutational load and the predicted neo-antigen load have been shown to broadly correlate with outcomes to ICPB in multiple studies.1–3 By utilizing the dual-tumor model, we were able to

**Figure 5.** Strong dLN neo-antigen responses to UNC45a predict combination ICPB outcomes.

a. Schema of treatment and surgery. b. Bilateral tumor growth curve over time. Arrows represent surgery and treatment time (n = 22). Responses to therapy are depicted by black solid lines. Pre-treatment reactivity to c. mutant UQCRC2, and d. mutant UNC45a in the dLN of responders (n = 12) and non-responders (n = 10). Student’s T-test; error bars mean ± SD.
Figure 6. Boosting UNC45a specific T cells does not improve responses to combination ICPB.

a. Schema of tumor inoculation, vaccination, surgery and treatment. b. Post-vaccination IFNγ production in response to neo-antigen peptides. Tumor growth curves of c. control, d. Montanide vehicle control + ICPB, e. ICPB only and f. UNC45a vaccine + ICPB groups. g. Survival plot and log-rank (Mantel Cox). h. IFNγ responses to UNC45a peptide in responders and non-responders to ICPB. Unpaired Student’s T-test.
show that the variation in neo-antigen response can be predictive of subsequent ICPB outcomes. We observed that strong dLN neo-antigen response is often accompanied by a higher background response to media and a strong response to other antigens. Overall it remains unclear if these dLN neo-antigen responses participate, or are even relevant, for a successful anti-tumor response i.e. the immune responses against neo-antigens in the dLN could potentially be secondary to events occurring in the tumor. Lesterhuis et al. showed that by day 6, identical tumors in genetically identical mice exhibit stochastically heterogeneous networks of gene expression that correlate with ICPB outcome. These responses may be a surrogate of a ‘hot’ tumor or a more active immune system. One argument that the latter notion is not the whole story is the observation that the neo-antigen response that best correlated with outcome was the one seen after ICPB – if the finding was entirely due to ‘hot tumors’, all neo-antigen reactivities would have been similarly boosted. Regardless, our observations have potential clinical significance. If the presence of a strong neo-antigen response in the dLN predicts response to ICPB therapy, this could be useful as a predictive biomarker in patients with accessible tumor dLN. Once relevant reactivities are identified in lymph nodes, they may then be able to be tracked in blood. Also, the finding that the dLN is the major site cross-priming neo-antigen specific T-cells processes means that local interventions in the region of the dLN may alter systemic anti-tumor immunity sufficient to make them a targetable site to boost anti-tumor response. Given this, it was logical to ask if ICPB unresponsiveness could be reversed if the low UNC45a response in some animals was boosted by pre-treatment vaccination. Interestingly, although an UNC45a vaccine boosted UNC45a specific responses, it failed to improve ICPB outcome. It is unclear if this is simply due to an inadequate vaccination response compared to other more aggressive vaccination strategies or that UNC45a is not a target for tumor rejection or that such responses are just a surrogate of the host’s immune responsiveness. It is also possible that a single antigen might be insufficient to induce a complete response alone, as it has been shown that vaccination with two neo-antigens induces optimal tumor rejection compared to either neo-antigen alone.

Selection of neo-antigen candidates to screen for immune responses is crucial. In our study, our strongly immunogenic neo-antigen UNC45a minimal epitope was not the highest predicted MHC-I binder. This was also evident in other studies where minimal epitopes with intermediate predicted MHC-I binding affinities were immunogenic. Furthermore, UQCRC2 and UNC45a peptide sequences (Supplementary figure 1) did not present any distinct immunological features, such as predicted binding affinity or predicted immunogenicity score. This highlights the challenges in neo-antigen discovery, epitope prediction, and the inherent disadvantage of relying on in silico predictions alone. A potential strategy to partially overcome this is the use long peptides spanning all the possible minimal epitopes. The predictive value of functional, neo-antigen responses to ICPB outcomes in our study provides the impetus to further examine these neo-antigens as vaccination targets. Importantly, neo-antigen T cell responses could be a potential biomarker in the clinic for ICPB responses whilst neo-antigen vaccination could be used to either treat or to ‘prime’ patients for a potential ICPB response. Further studies in the combination of neo-antigen vaccination strategies are required to provide better insights on how this can be achieved.

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References

1. Rizvi NA, Hellmann MD, Snyder A, Kvistborg P, Makarov V, Havel JF, et al. Mutational landscape determines sensitivity to PD-1 blockade in non-small cell lung cancer. Science. 2015;348(6230):124–128. doi:10.1126/science.aaa1348.
2. Snyder A, Makarov V, Merghoub T, Yuan J, Zaretsky JM, Desrichard A. Genetic basis for clinical response to CTLA-4 blockade in melanoma. N Engl J Med. 2014;371(23):2189–2199. doi:10.1056/NEJMoa1406498.
3. Van Allen EM, Miao D, Schilling B, Shukla SA, Blank C, Zimmerman L, Sucker A, Hillen U, Geukes Foppen MH, Goldinger SM, et al. Genomic correlates of response to CTLA-4 blockade in metastatic melanoma. Science. 2015;350(6257):207–211. doi:10.1126/science.aad0095.
4. Carreno BM, Magrini V, Becker-Hapak M, Kaabinejad S, Hundal J, Petti AA, Ly A, Lie W-R, Hildebrand WH, Mardis ER, et al. Cancer immunotherapy. A dendritic cell vaccine increases the breadth and diversity of melanoma neoantigen-specific T cells. Science. 2015;348(6236):803–808. doi:10.1126/science.aab3828.
5. Ott PA, Hu Z, Keskin DB, Shukla SA, Sun J, Bozym DJ, et al. An immunogenic personal neoantigen vaccine for patients with melanoma. Nature. 2017;547(7662):217–221. doi:10.1038/nature23003.
6. Sahin U, Derhovanessian E, Miller M, Klocy BP, Simon P, Lower M, Bukur V, Tadmor AD, Luxemburger U, Schrörs B, et al. Personalized RNA mutanome vaccines mobilize poly-specific therapeutic immunity against cancer. Nature. 2017;547(7662):222–226. doi:10.1038/nature23003.
7. Stronen E, Toebes M, Kelderman S, van Buuren MM, Yang W, van Rooyen J, Donia M, Bosch M-L, Lund-Johansen F, Olweus J, et al. Targeting of cancer neoantigens with donor-derived T cell receptor
38. Yadav M, Jhunjhunwala S, Phung QT, Lupardus P, Tanguay J, Bumbaca S, Franci C, Cheung TK, Fritsche J, Weinschenk T, et al. Predicting immunogenic tumour mutations by combining mass spectrometry and exome sequencing. Nature. 2014;515(7528):572–576. doi:10.1038/nature14001.

39. Castle JC, Kreiter S, Diekmann J, Lower M, van de Roemer N, de Graaf J, Selmi A, Diken M, Boegel S, Paret C, et al. Exploiting the mutanome for tumor vaccination. Cancer Res. 2012;72(5):1081–1091. doi:10.1158/0008-5472.CAN-11-3722.

40. Duan F, Duitama J, Al Seesi S, Ayres CM, Corcelli SA, Pawashe AP, Blanchard T, McMahon D, Sidney J, Sette A, et al. Genomic and bioinformatic profiling of mutational neoepitopes reveals new rules to predict anticancer immunogenicity. J Exp Med. 2014;211(11):2231–2248. doi:10.1084/jem.20141308.

41. Wei SC, Duffy CR, Allison JP. Fundamental mechanisms of immune checkpoint blockade therapy. Cancer Discov. 2018;8(9):1069–1086. doi:10.1158/2159-8290.CD-18-0367.

42. Hufeldt MR, Nielsen DS, Vogensen FK, Midtvedt T, Hansen AK. Variation in the gut microbiota of laboratory mice is related to both genetic and environmental factors. Comparative Med. 2010;60:336–342.

43. Marzo AL, Lake RA, Lo D, Sherman L, McWilliam A, Nelson D, Robinson BW, Scott B. Tumor antigens are constitutively presented in the draining lymph nodes. J Immunol. 1999;162(10):5838–5845.

44. Jackaman C, Lansley S, Allan JE, Robinson BWS, Nelson DJ. IL-2/CD40-driven NK cells install and maintain potency in the anti-mesothelioma effector/memory phase. Int Immunol. 2012;24(6):357–368. doi:10.1093/intimm/dxs005.