NLRC5/CITA expression correlates with efficient response to checkpoint blockade immunotherapy

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Checkpoint blockade-mediated immunotherapy is emerging as an effective treatment modality for multiple cancer types. However, cancer cells frequently evade the immune system, compromising the effectiveness of immunotherapy. It is crucial to develop screening methods to identify the patients who would most benefit from these therapies because of the risk of the side effects and the high cost of treatment. Here we show that expression of the MHC class I transactivator (CITA), NLRC5, is important for efficient responses to anti-CTLA-4 and anti-PD1 checkpoint blockade therapies. Melanoma tumors derived from patients responding to immunotherapy exhibited significantly higher expression of NLRC5 and MHC class I-related genes compared to non-responding patients. In addition, multivariate analysis that included the number of tumor-associated non-synonymous mutations, predicted neo-antigen load and PD-L2 expression was capable of further stratifying responders and non-responders to anti-CTLA4 therapy. Moreover, expression or methylation of NLRC5 together with total somatic mutation number were significantly correlated with increased patient survival. These results suggest that NLRC5 tumor expression, alone or together with tumor mutation load constitutes a valuable predictive biomarker for both prognosis and response to anti-CTLA-4 and potentially anti-PD1 blockade immunotherapy in melanoma patients.

Checkpoint blockade immunotherapy has emerged as one of the most promising strategies to treat patients with various cancers. Although many successes in single or combinatory use of anti-CTLA-4, PD-1 or PD-L1/2 antibodies have been documented in a variety of malignancies, responses are only typically observed in a minority of patients for any given regimen. Considering the substantial risk of autoimmune side effects and the high cost of the treatment, it is critical to develop screening methods to identify the subsets of patients who would most benefit from these therapeutics. While immunohistochemistry for PD-1/PD-L1 has been approved by the FDA, neo-antigen load, copy number alterations, TCR sequencing, multi-parameter flow cytometry, SERPINB mutations or nCounter gene expression profiling have been assessed for predicting responses to checkpoint blockade therapies, their predictive power and usefulness as potential biomarkers are limited.

Successful cancer growth and progression relies on the combination of both suppression and evasion of the host immune system. One of the major mechanisms of immune suppression in the tumor microenvironment is through impaired effector T cell function and can occur through multiple mechanisms such as increased expression of inhibitory immune checkpoint molecules, immunosuppressive enzymes and recruitment of immunosuppressive immune cells. Similarly, immune evasion from CD8+ T cells through defects in the MHC class I-mediated antigen presentation pathway is a common occurrence, with MHC class I loss or reduction reported in lung cancer (93%), prostate cancer (91%), cervical cancer (90%), pancreas cancer (86%), breast cancer (84%), colorectal cancer (78%), among others. Impaired MHC class I may manifest through many ways.
mechanisms such as loss of heterozygosity, gene mutations, epigenetic suppression or most importantly, down-regulation at the transcriptional level\(^1\). Indeed it has recently been shown that the MHC class I transactivator (CITA), \(\text{NLRC5}^{2,23}\), is a major suppression target to facilitate immune evasion in multiple cancers\(^2\). CITA/NLRC5 is an IFN-\(\gamma\)-inducible nuclear protein\(^{25-27}\) and transcriptionally regulates MHC class I gene activation via a CITA enhancer-like complex that specifically associates with MHC class I gene promoters\(^{26,29}\). As a master transcriptional activator of the MHC class I antigen presentation pathway, CITA/NLRC5-dependent MHC class I genes include \(\text{HLA}-A,-B,-C,-E,-\text{F}\) as well as the immunoproteasome component \(\text{LMP2 (PSMB9)}\), peptide transporter \(\text{TAP1}\) and \(\beta2\)-microglobulin (\(\text{B2M}\))\(^{23,26,29,30}\). The functional consequences of \(\text{NLRC5}\) deficiency in vivo constitutes failure of effective immune responses, accompanied with increased susceptibility to pathogen infection due to reduced constitutive and inducible expression of MHC class I gene expression and subsequently poor CD8\(^+\) T cell activation\(^{31-35}\). On the other hand, overexpression of \(\text{NLRC5}\) in cancer models has demonstrated improved tumor immunogenicity\(^3\). In human cancers, genetic and epigenetic changes in \(\text{NLRC5}\) gene are associated with impaired expression of MHC class I and related genes and reduced activity of CD8\(^+\) cytotoxic T cells\(^{24}\). Strikingly, increased expression of \(\text{NLRC5}\) is highly associated with improved 5-year survival of patients with skin, rectal, bladder, uterine, cervical or head/neck cancers, thus showing \(\text{NLRC5}\) expression is potentially valuable as a prognostic marker\(^3\).

**Results**

The expression of \(\text{NLRC5}\) and MHC class I associated genes are correlated with response to anti-CTLA-4 antibody therapy.

Since \(\text{NLRC5}\) expression is required for efficient cytotoxic CD8\(^+\) T cell responses, we hypothesized that \(\text{NLRC5}\) may be important for mediating the clinical benefits of cancer patients treated with checkpoint inhibitors\(^{24,29}\). We analyzed and compared the gene expression level of \(\text{NLRC5}\) and its dependent genes in melanoma between the groups who benefitted from the treatment (responder) and who did not benefit (non-responder). Among the melanoma patient cohort who received anti-CTLA-4 checkpoint blockade therapy, we observed a reduction in the gene expression level of \(\text{NLRC5}\)-dependent MHC class I and CD8\(^+\) T cell genes in non-responders versus responders (Fig. 1a). Gene set enrichment analysis indicated that this \(\text{NLRC5}\)-dependent MHC class I and CD8\(^+\) T cell gene set was upregulated in responders (Fig. 1b). Among these, we found that \(\text{NLRC5}\) expression was significantly elevated in the group who benefitted from the anti-CTLA-4 therapy (Fig. 1c). Because of the role of \(\text{NLRC5}\) as a major regulator of MHC class I and related genes, the expression of \(\text{NLRC5}\) is correlated with \(\text{HLA}-B\) and \(\text{B2M}\) in various cancers\(^{24,37}\) as well as in this melanoma patient cohort (Fig. S1). In addition to \(\text{NLRC5}\), the responder group exhibited higher expression of \(\text{HLA}-B\) than the non-responder group, and \(\text{B2M}\) showed a similar trend although it was not statistically significant with this cohort size (Fig. 1d). \(\text{NLRC5}\) is required for optimal recruitment and activation of CD8\(^+\) T cells in cancers\(^{24,37}\). As expected, the expression of \(\text{NLRC5}\) in various cancers\(^{24}\) and in this melanoma cohort was also correlated with the expression level of markers for CD8\(^+\) T cell activation, \(\text{CD8A}\) and granzyme A (\(\text{GZMA}\))/perforin (\(\text{PRF1}\)), but not \(\text{CD56}\), a marker for NK cells (Fig. S1). The responder group exhibited higher expression of \(\text{GZMA}\) and \(\text{PRF1}\) (Fig. 1e). Although \(\text{GZMA}\) and \(\text{PRF1}\) are expressed in both CD8\(^+\) T cells and NK cells, the high expression of \(\text{GZMA}\) and \(\text{PRF1}\) was likely due to activated CD8\(^+\) T cells rather than NK cells, since \(\text{CD56}\) expression in the responder group was not significantly different than that of the non-responder group (Fig. 1e). These data suggest that \(\text{NLRC5}\) and \(\text{NLRC5}\)-mediated MHC class I dependent CD8\(^+\) T cell activation is important for effective response to anti-CTLA-4 checkpoint immunotherapy.

**\(\text{NLRC5}\) expression and load of mutation or neoantigen provide predictive information for the response to anti-CTLA-4 therapy.** It has been shown recently that neoantigen load is an important predictor of responses to anti-CTLA-4 therapy; high neoantigen loads in melanoma are correlated with favorable responses to anti-CTLA-4 therapy\(^3\). Similarly, the number of total mutations (mutation load), which has been demonstrated to be highly correlated with neoantigen load in various cancers as well as samples in this cohort (Fig. S2), is important in predicting response\(^{6,39}\). In order to test if the addition of mutation/neoantigen load to \(\text{NLRC5}\) expression would improve predictions, we performed multivariate analysis by logistic regression treating these variables as covariates. Consistent with a previous report\(^{38}\), responding patients in this study also showed higher neoantigen load and number of tumor-associated mutations (Fig. 2a). Scatter plots for \(\text{NLRC5}\) expression combined with neoantigen load or mutation number showed non-responder groups were clearly separated from responders (Fig. 2b). Patients were then stratified by \(\text{NLRC5}\) expression and neoantigen load or number of mutations, yielding four groups (high/high, high/low, low/high, and low/low). The response rate in the group with low \(\text{NLRC5}\) expression and low neoantigen load (or low mutation number) was significantly less than that of the group with high \(\text{NLRC5}\) expression and high neoantigen load (or high mutation number) (Fig. 2c). These results suggest that two variables, \(\text{NLRC5}\) expression and neoantigen load (or mutation number) may be useful to jointly identify non-responders. ROC analysis based on the prediction equation from logistic regression showed a substantial increase in the area under the curve (AUC) when mutation/neoantigen load was included as a predictor in the regression model (Fig. 2d). For the model that included \(\text{NLRC5}\) and mutation load, 100% sensitivity was obtained at a 46\% false positive rate (Fig. 2d, left). Without mutation load, a false positive rate of 91\% was required to achieve 100% sensitivity. Similarly, false positive rate with 100% sensitivity was improved to 64\% when neoantigen load was included as a predictor (Fig. 2d, right). These data further indicate that analysis with two variables are useful to predict the patient population who will not respond to anti-CTLA-4 therapy.

**Combination of PD-L2 expression with \(\text{NLRC5}\) expression and mutation or neoantigen load are sensitive predictors for responses to anti-CTLA-4 therapy.** Based on the superior performance of our ROC curve analyses using \(\text{NLRC5}\) and mutation or neoantigen load (Fig. 2d), we sought to further...
improve prediction by adding more variables. We reasoned that CTLA-4, PD-1, PD-L1 or PD-L2 might be good candidates because their expression has been proposed to be correlated with responses to checkpoint blockade immunotherapies\(^38,40,41\); therefore, we first examined the correlation between NLRC5 expression and the expression of these genes. The expression of NLRC5 exhibited intermediate to high correlation with the expression of CTLA-4 (Pearson's correlation coefficient 0.70) and PD-1 (0.83), while the correlation between NLRC5 expression and expression of PD-L1 (0.44) and PD-L2 (0.54) was lower (Fig. S3), suggesting that CTLA-4 and PD-1 might not be good predictive variables to partner with NLRC5. ROC curve analyses revealed that the AUC was the highest when NLRC5 expression, mutation load and PD-L2 expression were included as covariates (Table S1).

![Figure 1](https://example.com/figure1.png)

**Figure 1.** The expression of NLRC5-dependent MHC class I and CD8+ T cell genes are correlated with response to anti-CTLA-4 antibody therapy. Patients groups who benefitted from anti-CTLA4 antibody therapy (Response, n = 14) and who did not (Nonresponse, n = 23) were analyzed for differential gene set enrichment by (a) heatmap and (b) GSEA as well as individual gene expression levels of (c) NLRC5, (d) HLA-B, B2M, (e) CD8A, granzyme A (GZMA), perforin (PRF1) and CD56. Bar represents the median value. P-values calculated using Mann–Whitney U test. NES, normalized enrichment score.
**a**

Mutations or neoantigen load (log10)

- **Response** (n=13) - Red dots
- **Nonresponse** (n=22) - Blue dots

**b**

- **Response** (n=13)
- **Nonresponse** (n=22)

- **p = 0.043**
- **p = 0.052**

Mutation load (log10) vs. NLRC5 expression (log2 normalized count)

**c**

Response rate (%)

- **Response**
- **Nonresponse**

NLRC5 Mutation:
- H, H, L, H, L

- **p = 0.032**

NLRC5 Neoantigen:
- H, H, L, H, L

- **p = 0.051**

**d**

- **NLRC5** (AUC = 0.69 ± 0.098)
- **Mutation load** (AUC = 0.70 ± 0.091)
- **NLRC5 + Mutation load** (AUC = 0.74 ± 0.085)
- **Neoantigen load** (AUC = 0.67 ± 0.093)
- **NLRC5 + Neoantigen load** (AUC = 0.71 ± 0.091)

Sensitivity (%) vs. 100-Specificity (%)
methylation group respectively (HR = 0.45 and HR = 2.31) (Fig. 4a), NLRC5 expression group and high NLRC5 expression and low NLRC5 methylation showed significantly better prognosis than the low NLRC5 expression/mutation or neoantigen load. The response rate (%) to the therapy among each group was calculated. Patients carrying higher value of the median are defined as high group (H), those carrying lower value of the median are defined as low group (L) in respective variables. Statistical significance between the groups of high NLRC5 expression/high mutation or neoantigen load and low NLRC5 expression/low mutation or neoantigen load were determined by the $\chi^2$ test. (d) ROC curves for logistic regression models using the respective combination of NLRC5 expression, mutation load and neoantigen load. The numbers with arrow are showing false positive rate with 100% sensitivity. AUC (area under the curve) ± SE (standard error) is depicted.

**Combination of NLRC5 expression and load of mutation or neoantigen provide prognostic information for the response to anti-CTLA-4 therapy.** Previously it was shown that NLRC5 expression is correlated with prognosis of patients with multiple cancer types\(^4\). The multivariable logistic regression including NLRC5 expression together with mutation load or neoantigen load indicated that the analysis of two variables would be superior to predict responses to anti-CTLA-4 checkpoint blockade therapy (Fig. 2b–d). Since these variables are critical for immune surveillance against cancer, we hypothesized that an association would be observed with patient prognosis and overall survival. Using melanoma patient data from the TCGA database, we performed a survival curve analysis using Kaplan–Meier estimates for overall survival and a multivariate Cox proportional hazards model for hazard ratios. The cohort was divided into two groups based on values higher or lower than the median for mutation load, NLRC5 expression and NLRC5 promoter methylation. The high mutation patient group demonstrated better prognosis than the low mutation group (HR = 0.44) (Fig. 4a). The groups with high NLRC5 expression and low NLRC5 methylation showed significantly better prognosis than the low NLRC5 expression group and high NLRC5 methylation group respectively (HR = 0.45 and HR = 2.31) (Fig. 4a), supporting previous reports that NLRC5 expression and methylation is correlated with prognosis of melanoma patients\(^4\). Survival curve analysis of four groups stratified by the level of NLRC5 expression and mutation load demonstrated that the high NLRC5 expression/high mutation load group showed better prognosis than the low NLRC5 expression/low mutation load group (Fig. 4b). Similarly, survival curve analysis for four groups stratified by the level of NLRC5 promoter methylation and mutation load showed that NLRC5 methylation high/mutation low group is a high risk group with poor prognosis, and the NLRC5 methylation low/mutation high group is a lower risk group with better prognosis (Fig. 4b). Taken together, these data indicate that multivariate analysis using NLRC5 expression/methylation status with mutation load is superior to single variable analysis and may be of value as a prognostic biomarkers in melanoma.

**The expression of NLRC5 and MHC class I-associated genes are correlated with response to anti-PD1 antibody therapy.** Based on our observations of the potential value of NLRC5 as a biomarker in the anti-CTLA4 treatment cohort, we were inspired to extend the analysis to melanoma patient cohorts treated with anti-PD1 therapy. Gene set enrichment analysis indicated that the NLRC5-dependent MHC class I and CD8+ T cell gene set was also shown to be reduced in patients who did not respond to anti-PD1 therapy (Fig. 5a,b). Similar to anti-CTLA4-treated patient cohort, NLRC5 and HLA-B was reduced in non-responders (Fig. 5c), along with a similar trend for B2M in anti-PD1-treated melanoma patients (Fig. 5d). Similarly, CD8+ T cell markers, CD8A, PRF1 and GZMA were decreased with no change to CD56 (Fig. 5e). The predictive value of NLRC5 expression alone in the anti-PD1 cohort was comparable (AUC = 0.71) to what was observed in anti-CTLA4 (Fig. 5f) and NLRC5 expression clearly stratified patients into a high and low overall survival group upon Kaplan-Meier analysis (Fig. 5g). These data indicate that NLRC5 expression level may also be important for effective response to anti-PD1 monotherapy and may provide predictive information.

**Discussion**

Discovery of inhibitory receptors on T cells and development of monoclonal antibodies against them has led to widespread usage of checkpoint blockade therapy in various cancers\(^7\). Although these therapies are effective for many cancer patients, complete response rate ranges from around 20% for anti-CTLA-4 antibody therapy\(^2,43\) to 30% for anti-PD/anti-PD-L1 therapy in the case of melanoma\(^41,43\). These treatments are quite expensive and when ineffective create a significant financial burden on patients and the health care system\(^44\). Although it is known that the expression of PD-1/PD-L1, mutation and neoantigen load correlate with responses checkpoint blockade therapy, their predictive power is low, resulting in the treatment of many patients for whom these therapies are ineffective.

This study suggests that NLRC5 is a biomarker to predict the outcome of CTLA-4 blockade therapy. Since checkpoint therapy relies on T cell activation and NLRC5 is critical for MHC class I-dependent cytotoxic T cell activation, it is not unexpected that NLRC5 may play an important role in the response to checkpoint blockade therapy. Indeed, reduced MHC class I immunostaining and gene expression in pretreatment biopsies from...
anti-CTLA-4 treated melanoma patients predicts resistance to therapy and reduced survival. Similarly, others have independently shown that increased immunoproteasome expression (PSMB8, PSMB9) is correlated with better response to anti-CTLA-4 or anti-PD-1 therapy. Although NLRC5 expression alone has relatively weak predictive power (Fig. 2d), in combination with other variables it yields improved predictive performance. In particular, NLRC5 expression and neoantigen load/mutation number exhibited a low degree of multi-collinearity and are weakly correlated (Pearson’s coefficient 0.3 and 0.27, respectively, Fig. S3). Combining NLRC5 expression

Figure 3. Combination of PD-L2 expression with NLRC5 expression and mutation or neoantigen load are sensitive predictors for responses to anti-CTLA-4 therapy. (a) Scatterplots for NLRC5 and PD-L2 expression with mutation load (left panel) or neoantigen load (right panel) for response (n = 13) and nonresponse (n = 22) groups. (b) ROC curves for logistic regression models using the respective combination of PD-L2 expression, NLRC5 expression, mutation load and neoantigen load. The numbers with arrow are showing false positive rate with 100% sensitivity. AUC (area under the curve) ± SE (standard error) is depicted.
and mutation numbers demonstrated better AUC values and a lower false positive rate at 100% sensitivity (Fig. 2d). These data indicate that the combination of \( \textit{NLRC5} \)/mutation load is superior to these variables alone in identifying non-responders. In contrast to the low correlation between \( \textit{NLRC5} \) expression and mutation load, the expression of \( \textit{CTLA-4} \), \( \textit{PD-1} \), \( \textit{PD-L1} \) or \( \textit{PD-L2} \) relative to \( \textit{NLRC5} \) expression carry high to intermediate correlation (Pearson's coefficient 0.70, 0.83, 0.44 or 0.54, Fig. S3). It appears that \( \textit{PD-L2} \) is the best variable to combine with \( \textit{NLRC5} \) and mutation number for purposes of prediction (Table S1) and is in line with \( \textit{PD-L2} \) being a predictive marker in anti-PD-1 checkpoint therapy47. Future discovery of other variables will further

**Figure 4.** Combination of \( \textit{NLRC5} \) expression and load of mutation or neoantigen provide prognostic information. (a) Kaplan–Meier estimates of five year overall survival of patients with high and low mutation load (Left), \( \textit{NLRC5} \) gene expression (Middle), and \( \textit{NLRC5} \) methylation (Right). Patients in the TCGA melanoma cohort were stratified by medians into high and low groups (n = 159 and n = 160). (b) Kaplan–Meier estimates of five year overall survival of patients with varying levels of two factors, \( \textit{NLRC5} \) expression and mutation load (Left) and \( \textit{NLRC5} \) methylation and mutation load (Right). Patients were stratified by two factors (\( \textit{NLRC5} \) expression/\( \textit{NLRC5} \) methylation and mutation load) in a similar fashion with (a), yielding four groups (high \( \textit{NLRC5} \) expression/\( \textit{NLRC5} \) methylation and high mutation load, likewise, high and low, low and high, low and low). Pairwise log-rank test was used to analyze the survival in indicated pairs. Hazard ratio (HR) and 95% confidence interval (CI) was determined by multivariate analysis using Cox regression model (see Methods). *p < 0.05; **p < 0.01; ***p < 0.001.
Figure 5. The expression of NLRC5 and NLRC5-dependent MHC class I and CD8+ T cell genes as a predictor to anti-PD1 therapy. Patients groups who benefitted from anti-PD1 therapy (Response, n = 22) and who did not (Nonresponse, n = 19) were analyzed for differential gene set enrichment by (a) heatmap and (b) GSEA as well as individual gene expression levels of (c) NLRC5, (d) HLA-B, B2M, (e) CD8A, granzyme A (GZMA), perforin (PRF1) and CD56. Bar represents the median value. P-values calculated using Mann-Whitney U test. NES, normalized enrichment score. (f) ROC curve for logistic regression model using NLRC5 expression. The numbers with arrow are showing false positive rate with 100% sensitivity. AUC (area under the curve) ± SE (standard error) is depicted. (g) Kaplan–Meier estimates of five year overall survival of patients with high and low NLRC5 gene expression, stratified by median expression (n = 20 and n = 21). Hazard ratio (HR) and 95% confidence interval (CI) was determined by multivariate analysis using Cox regression model (see Methods). **p < 0.01.
increase predictive power for response to the checkpoint therapy in melanoma. Although this study involved only a relatively small number of melanoma patients who received anti-CTLA-4 checkpoint therapy, we demonstrated an almost identical relationship between NLRC5 and differentiation of non-responders from responders of anti-PD-1 checkpoint therapy, albeit mutation/neoantigen load data was not available for us to fully replicate our model (Fig. 5). Anti-PD-1/PD-L1 antibody therapy is based on similar mechanisms to increase anti-tumor immune responses. Thus, it is feasible that NLRC5 expression/mutation load might also be useful for predicting outcomes of other cancer patients treated with anti-PD-1/PD-L1 antibody therapy. Checkpoint blockade therapy was initially tested in melanoma patients, but has been expanded to a dozen cancer types including lung, breast and kidney. Therefore, investigations into the role of NLRC5 expression and mutation load for the prediction of treatment outcomes in these cancers is of interest.

In summary, this study identified the expression of NLRC5 as a novel predictive biomarker for immune checkpoint blockade therapies in melanoma. Multivariate analysis using NLRC5 seems to have significant potential as a predictor of patient response to checkpoint blockade therapy. Validation studies using a larger independent cohort is needed to corroborate these results and to further refine the predictive relationships for checkpoint blockade treatment outcomes.

**Materials and methods**

**Data sets.** *Clinical data for anti-CTLA4 therapy.* The cohort for the analysis of response to anti-CTLA-4 therapy (ipilimumab) was obtained through Database of Genotypes and Phenotypes (dbGaP)48,49, accession number phs000452.v2.p138.

TCGA. Data for survival analysis of melanoma was obtained through the Cancer Genome Atlas (TCGA) data portal (https://tcga-data.nci.nih.gov/tcga), Skin Cutaneous Melanoma (SKCM). Gene expression data (mRNASeq: illuminahiseq_rnaseqv2-RSEM_genes_normalized_data), DNA methylation data (humanmethyl-ation450-within_bioassay_data_set_function) and clinical data (Merge_Clinical) were accessed through GDAC Firehose (gdac.broadinstitute.org). Somatic mutation data were accessed through the cBioPortal50.

**Clinical data for anti-PD1 therapy.** The raw sequence files for the cohort of response to anti-PD1 therapy was obtained from the European Nucleotide Archive (ENA) under accession PRJEB23709 and matched to clinical data from source study32.

**Patient profiling.** *dbGap dataset (anti-CTLA4 therapy).* A patient population of 37 metastatic melanoma patients who had taken ipilimumab monotherapy was analyzed. Patients were stratified by clinical benefit status as described previously48. Response to ipilimumab was defined as CR (complete response) or PR (partial response) by Response Evaluation Criteria in Solid Tumors (RECIST) criteria or SD (stable disease) by RECIST criteria32 with overall survival greater than 1 year. Non-response to ipilimumab was defined as PD (progressive disease) or SD (stable disease) by RECIST criteria with overall survival less than 1 year.

**ENA dataset (anti-PD1 therapy).** A patient population of 41 melanoma patients treated with anti-PD1 therapy using either nivolumab (n = 9) or pembrolizumab (n = 32) with available pre-treatment sampled tumor RNA-seq data was analyzed. Patient stratification into responder and non-responders was performed under the same criteria used with the anti-CTLA-4 patients.

**Gene expression analysis.** RNA sequences were downloaded and converted to FastQ file format using SRA Toolkit v2.6.3. Paired-end reads were checked to trim for low quality bases and adaptor sequences using Trimomatic33. For the dbGap dataset, filtered reads were mapped to the GRCh37/hg19 assembly using TopHat v2.0.1334 and HTSeq55 was then used to generate raw read counts per gene using intersection-nonempty parameter to account for ambiguous read mappings. For the ENA dataset, filtered reads were mapped to the GRCh38/hg19 assembly using HISAT236 with GNU-parallel37 and BAM files indexed and sorted by SAMtools58. featureCounts59 was used to count reads mapped to each gene. Gene expression values were generated for further analyses using DESeq260, following recommended guidelines by the authors. The expression levels of NLRC5, HLA-B, B2M, CD8A, granzyme A (GZMA), perforin (PRF1) and CD56 measured by RNA sequencing (RNA-seq) were compared between responders and non-responders using the Mann–Whitney U Test.

**Gene set enrichment analysis (GSEA).** Gene Set Enrichment Analysis (GSEA, http://www.broad.mit.edu/gsea/)61 was used to assess differential expression of NLRC5-dependent MHC class I and CD8+ T cell gene set between response and non-response group to anti-CTLA4 or anti-PD1 therapy. The gene set tested was based on knowledge of the literature concerning NLRC5 and the MHC class I antigen presentation pathway32,34,37.

**Mutation and neoantigen analysis.** For analyses involving numbers of mutation and neoantigen (mutation load and neoantigen load, respectively), 35 of the 37 melanoma patients treated with ipilimumab had data available38. Those were compared between responders and non-responders to ipilimumab using the Mann–Whitney U Test. The bidimensional combination of NLRC5 expression and mutation or neoantigen load was assessed and p-values were calculated using Hotelling’s T2 Test to compare between responder and non-responder to ipilimumab. Next, to evaluate the influence of those variables for response to ipilimumab, cohort was divided into four groups based on the level of NLRC5 expression and mutation or neoantigen load and was calculated for the response rate to ipilimumab. Patients carrying higher value of the median are defined as...
high group, those carrying lower value of the median are defined as low group in respective variables. Statistical significance between the groups of high NLRC5 expression/high mutation or neoantigen load and low NLRC5 expression/low mutation or neoantigen load was determined by the χ² test.

Expression values for other genes known as predictors of response to anti-CTLA-4 therapies, such as CTLA-4, PD-1, PD-L1 and PD-L2 were combined with NLRC5 expression and mutation or neoantigen load and represented in a three-dimensional scatterplot.

**Logistic regression analysis.** Logistic regression models were fitted with different combinations of the following covariates: the values for expression of 5 genes (NLRC5, PD-L1, PD-L2, CTLA-4 and PD-1), mutation load, and neoantigen load. Up to 3 of these variables were considered for the regression models at a time. Samples with missing values were eliminated before fitting the regression. Multicollinearity was assessed through calculation of variance inflation factors and Pearson’s correlation coefficient (Fig. S3). A scatterplot matrix was created with fitted curves and regression lines and the distribution of each variable was inspected. An ROC curve was generated for each combination of covariates and AUC± standard error (SE) generated using the pROC package (version 1.8) in R. The training data was used as the prediction data for the ROC curves. Threshold values were determined at points where the sensitivity is 100%. These curves were plotted and a selection were reported. A bootstrapping procedure with 10,000 repetitions was used to estimate 95% confidence intervals for the curves as well as calculate a mean AUC. This was accomplished by sampling the cohort with replacement to create new groupings of data (the same size as the original cohort) then used to construct ROC curves. The AUC was calculated for each of these new curves. The confidence interval was determined by ordering the AUCs by value and returning the value at the index 2.5% of the length of the list away from the beginning and end. The best prediction model was chosen based on the highest mean AUC.

**Survival analysis.** Survival analysis was performed on a cohort of melanoma patients obtained from TCGA (n = 319) or ENA (n = 41). The TCGA cohort was filtered to only include patients who had had all three data types available (RNA-seq, mutation load and methylation) as well as clear cancer tumor stage record. NLRC5 methylation was assessed as the sum of the methylation of probe sites (cg08159663, cg07839457 and cg16411857) that negatively correlated with NLRC5 gene expression (Fig. S4). Survival curves were calculated using the Kaplan–Meier method depicting the difference in survival between the groups through division of the cohort into top and bottom 50% based on median NLRC5 expression, NLRC5 methylation or mutation load. Patients were stratified in a similar fashion by two variables (NLRC5 expression and mutation load) yielding four groups (high NLRC5 expression/high mutation load, high/low, low/high, low/low, respectively). The same was also performed for NLRC5 methylation and mutation load (high NLRC5 methylation/high mutation load, high/low, low/high, low/low, respectively). For univariate survival curves, statistical significance was assessed by log-rank test and for multivariate survival curves, pairwise log-rank test with Benjamini–Hochberg FDR correction was used. A Cox proportional hazard regression model was used to evaluate the effects of age and cancer stage as additional covariates. Cancer stage was not available for patients in the anti-PD-1 dataset.

**Statistical analysis.** Statistical analysis was performed using R 3.6.0 and RStudio 1.1.463.

**Ethical approval and ethical standards.** The data used in our study were obtained from public databases, therefore, ethical approval was not required.
citations

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Author contributions
S.Y., J.Y., G.L., V.E.J. and K.S.K. designed studies. S.Y., S.X.C., X.P., J.Y. and K.S.K. performed analysis. S.Y., S.X.C., J.Y. and K.S.K. wrote the paper.

Funding
This work was supported by grants from the American Lung Association LCD-507710, National Multiple Sclerosis Society, CSTR, TAM Genomics, Japan Society for the Promotion of Science (JSPS) Grant Number: 18H06135, SENSHIN Medical Research Foundation, Takeda Science Foundation to KSK, fellowship from Sumitomo Life Welfare and Culture Foundation to SY and JSPS Grant Number: 20K16271 to SXC.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary Information The online version contains supplementary material available at https://doi.org/10.1038/s41598-021-82729-9.

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