# Differential immune profiles distinguish the mutational subtypes of gastrointestinal stromal tumor

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Gastrointestinal stromal tumor (GIST) is the most common human sarcoma, frequently characterized by an oncogenic mutation in the *KIT* or *PDGFRA* gene. We performed RNA sequencing of 75 human GIST tumors from 75 patients, comprising what we believe to be the largest cohort of GISTs sequenced to date, in order to discover differences in the immune infiltrates of *KIT*- and *PDGFRA*-mutant GIST. Through bioinformatics, immunohistochemistry, and flow cytometry, we found that in *PDGFRA*-mutant GISTs, immune cells were more numerous and had higher cytolytic activity than in *KIT*-mutant GISTs. *PDGFRA*-mutant GISTs expressed many chemokines, such as CXCL14, at a significantly higher level when compared with *KIT*-mutant GISTs and exhibited more diverse driver-derived neoepitope:HLA binding, both of which may contribute to *PDGFRA*-mutant GIST immunogenicity. Through machine learning, we generated gene expression-based immune profiles capable of differentiating *KIT*- and *PDGFRA*-mutant GISTs, and identified additional immune features of high-PD-1- and -PD-L1-expressing tumors across all GIST mutational subtypes, which may provide insight into immunotherapeutic opportunities and limitations in GIST.

# Introduction

Gastrointestinal stromal tumor (GIST) is the most common sarcoma (1), often originating in the stomach or the small intestine. GIST is a paradigm for targeted molecular therapy, since the majority of GISTs contain an activating mutation in exon 11 of the *KIT* proto-oncogene, for which there are effective and well-tolerated tyrosine kinase inhibitors (2, 3). Imatinib mesylate, which inhibits the KIT oncoprotein, improved median survival in patients with advanced and metastatic GIST from 1 to 5 years (3), while adjuvant imatinib improved recurrence-free survival in patients with resectable disease (4).

*PDGFRA*-mutant GIST is the second most common form of GIST. Approximately half of *PDGFRA*-mutant GISTs contain a *D842V* substitution, which is inherently resistant to imatinib therapy (5). The majority of *PDGFRA*-mutant GISTs develop in the stomach (6, 7), while *KIT*-mutant GISTs can arise in the stomach, small intestine, or rectum (8), suggesting an innate and important biologic difference between these driver mutations. Other GIST mutation types also associate with predictable clinicopathologic features. For example, *KIT* exon 9-mutant GISTs nearly always develop in the small intestine (9); neurofibromin 1-mutant (*NF1*-mutant) GISTs are more commonly found in the duodenojejunal flexure (10); and succinate dehydrogenase-deficient

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(SDH-deficient) GISTs are indolent, multifocal, and among the few GISTs — along with those driven by kinase fusions — that metastasize via the lymphatic system (11, 12). The underlying biologic mechanisms linking mutation type and clinicopathologic features are not well understood.

Although *PDGFRA D842V* (*D842V*)-mutant GISTs do not respond to imatinib, natural history shows that recurrence-free survival is more favorable in patients with any *PDGFRA* mutation when compared with a *KIT* mutation (4). This finding, along with the predictable behavior of GIST mutation types, had us hypothesize that the mutational driver may impact other aspects of tumor biology, specifically the tumor microenvironment and host immune response. Our group has extensively characterized the immune infiltrate in *Kit*-mutant GIST using a genetically engineered mouse model and human specimens (13–17), providing a clear rationale for multiple immunotherapy trials in patients with GIST, but the immune response to *PDGFRA*-mutant GIST is currently unknown.

In this study, we performed RNA-Seq on 75 surgical specimens from 75 patients with GIST to characterize the immune infiltrates of different GIST mutations. Surprisingly, we discovered that *PDGFRA*-mutant GIST harbors more immune cells than *KIT*-mutant GIST; this may be related to oncogene-specific cytokine production or more HLA-diverse neoepitope recognition, suggesting that patients with *PDGFRA*-mutant GIST might benefit from therapeutic immunomodulation. We trained a random forest machine learning algorithm on RNA-Seq data from all of our *KIT*- and *PDGFRA*-mutant GISTs, as well as on a subset of untreated, primary, gastric (UPG) *KIT*- and *PDGFRA*-mutant GISTs, in order to characterize a *PDGFRA*-specific immune land-

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**Figure 1. ssGSEA identifies immune cell pathway enrichment in PDCFRA-mutant GIST.** ssGSEA of 75 GIST specimens, organized by mutational driver and increasing ESTIMATE score. Unsupervised row clustering grouped gene sets into 3 major categories based on cell cycle pathways, metabolic pathways, and immune pathways. Clinicopathologic characteristics of the 75 GIST specimens are shown in the annotation and in Supplemental Table 1.

scape. Finally, we identified the top immune features correlating with high PD-1 and PD-L1 expression across all GIST specimens to identify potential barriers to and synergistic opportunities for successful immune checkpoint blockade in GIST.

## Results

Single-sample gene set enrichment analysis identifies immune cell pathway enrichment in PDGFRA-mutant GIST. We performed RNA-Seq and principal component analysis (PCA) of 75 human GIST specimens comprising various mutation types (n = 37 KIT-mutant, n = 24 PDGFRA-mutant, n = 7 SDH-deficient, n = 4 multiple drivers, n = 2 WT, and n = 1 NFI-mutant; Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/JCI124108DS1). The combined PCA of all mutational drivers demonstrated clustering by mutational subtype (Supplemental Figure 1, top), which was most apparent in

*PDGFRA*-mutant and *SDH*-deficient GISTs (Supplemental Figure 1, bottom). Notably, *KIT*-mutant GISTs appeared to cluster into 3 distinct groups, which upon closer inspection of clinicopathologic features was related to treatment status and tumor location.

Given what our group has discovered regarding the immune infiltrate of GIST (13–15, 17), the metabolic characteristics of imatinib-treated GIST (18), and the association of cell cycle activity with GIST aggressiveness (19, 20), we used our sequencing data to perform single-sample gene set enrichment analyses (ssGSEAs) focused on immune, metabolic, and cell cycle pathways (Figure 1). Using published gene sets (21), along with assessment of previously published metabolic, cell cycle, and immune pathways (22), we identified 136 gene sets for inclusion in ssGSEA (Supplemental Table 2).

ESTIMATE and CYT scoring, which infer the quantity and cytolytic activity of the immune infiltrates, respectively (23, 24), revealed that *KIT*-mutant GIST had a range of immune cell

infiltration and cytolytic activity (Figure 1). Meanwhile, *SDH*deficient, *NF1*-mutant, *D842V*-mutant GISTs with a concurrent *CDKN2A* (p16) deletion, and WT GISTs (defined as non-*KIT*, non-*PDGFRA*, non-RAS-activated, and non-*SDH*-deficient, as previously described; ref. 25) exhibited generally low ESTIMATE and CYT scores and a lack of immune cell pathway enrichment.

Conversely, ESTIMATE, CYT, and ssGSEA identified increased immune cell infiltration, greater immune cell activity, and a significant enrichment of immune-related gene sets in *PDGFRA*-mutant GISTs — an unexpected finding that suggested that *PDGFRA*-mutant GISTs were more immunogenic than other GIST mutations (Figure 1). Cell cycle activity pathways did not clearly correlate with immune cell infiltration across all GIST driver mutation types, while metabolic activity appeared to inversely correlate with the quantity of immune infiltrate, specifically in *KIT* exon 11-mutant GISTs. Overall, these data show that while tumors driven by a particular GIST mutation can have variable immune profiles, *PDGFRA*-mutant GIST contains the strongest gene expression-based immune signature when compared with other GIST mutations.

PDGFRA-mutant GIST is more immunologically active than KIT-mutant GIST. In order to validate our ssGSEA finding that PDGFRA-mutant GIST was highly enriched in immune cell quantity and cytolytic activity, we first compared the clinicopathologic characteristics of all KIT- and PDGFRA-mutant GISTs (Supplemental Table 3; n = 61). Though not associated with immune cell infiltration in GIST, tumor size, mitotic rate, and tumor location are predictive of GIST aggressiveness and recurrence-free survival after surgical resection (26). PDGFRA-mutant GISTs were more commonly found in stomach when compared with KIT-mutant GISTs, and there were significantly more males in our PDGFRA-mutant GIST cohort. There was also a trend toward decreased mitotic activity in PDGFRA-mutant when compared with KIT-mutant GISTs (Supplemental Table 3). ESTIMATE and CYT scores, however, did not correlate with mitotic rate across all KIT- and PDGFRA-mutant GISTs (Supplemental Figure 2A). CYT score and the overall number of CD45 mRNA transcripts were significantly higher in PDGFRA-mutant when compared with KIT-mutant GISTs (Figure 2A), while there was also a trend toward increased ESTIMATE score and CD8 mRNA expression, suggesting that PDGFRA-mutant GISTs may exhibit greater immune cell infiltration and activation than KIT-mutant GISTs.

Since we observed significant differences in tumor location between *KIT*- and *PDGFRA*-mutant GISTs, and metastatic lesions and treatment status have also been shown to alter the immune infiltrate in GIST (14, 27), we then compared only UPG *PDGFRA*and *KIT*-mutant GISTs to minimize confounders and maximize the potential to observe important biologic differences that may be related to the oncogenic driver (Supplemental Table 4; n = 22). UPG *PDGFRA*- and *KIT*-mutant GISTs with a low mitotic rate exhibited significantly higher ESTIMATE scores when compared with UPG GISTs with a high mitotic rate (Supplemental Figure 2B), suggesting that immune cell infiltration may be partly related to GIST aggressiveness when controlling for the mutational driver in addition to tumor location and treatment status, mitotic rate did not inversely correlate with immune cell infiltration (Supplemental Figure 2C), implying that the oncogenic driver may also be contributing to differences in immune response. In fact, ESTIMATE score, CYT score, and the mRNA expression of CD45 were significantly greater in UPG *PDGFRA*-mutant GISTs when compared with UPG *KIT*-mutant GISTs, and expression of CD8 mRNA was also increased (Figure 2B). Furthermore, gene set enrichment analyses (GSEAs) showed that the overall immune response pathway, adaptive immune response pathway, antigen binding pathway, and  $\alpha/\beta$  T cell activation pathway (Figure 2C), as well as lymphocyte activation, lymphocyte differentiation, and B cell activity pathways (data not shown), were more significantly enriched in UPG *PDGFRA*-mutant compared with *KIT*-mutant GISTs, suggesting that the difference in immune infiltration and activity may be related to the oncogenic driver.

To further validate the observed differences in immune cell infiltrate between PDGFRA- and KIT-mutant GISTs, we performed IHC staining for CD45 and CD8 on KIT- and PDGFRA-mutant GISTs, including additional tumors not included in the sequencing cohort (Figure 2D). On gross microscopic examination, it appeared that PDGFRA-mutant GISTs contained more CD45<sup>+</sup> and CD8<sup>+</sup> cells, with a proportion of immune cells clustered around perivascular structures, a finding that we have found to be associated with adaptive immunity (our unpublished observations). Quantification of CD45 and CD8 IHC staining, including staining of the most infiltrated and cytolytically active KIT-mutant GIST in the KIT-mutant cohort (human GIST 203), showed that PDGFRA-mutant GISTs had significantly more CD45<sup>+</sup> and CD8<sup>+</sup> cells per ×20 high-power field (HPF) than KIT-mutant GISTs (Figure 2E). Moreover, flow cytometric analysis of KIT- and PDGFRA-mutant GIST specimens confirmed that PDGFRA-mutant GISTs harbored significantly more CD45<sup>+</sup> immune cells than KIT-mutant GISTs (Figure 2E). Together, these findings confirm that in PDGFRA-mutant GISTs, immune cells are more numerous and have higher cytolytic activity than in KIT-mutant GISTs with similar clinicopathologic features.

CIBERSORT and differential gene expression analysis identify unique immune signatures in GIST. To discover additional differences in the immune landscape between UPG PDGFRA- and UPG KIT-mutant GISTs, we profiled a previously defined set of 117 relevant immune features (Supplemental Table 5) (28, 29). In addition to ESTIMATE and CYT scores, this profile included scores derived from CIBERSORT, which infers various immune cell frequencies through RNA-Seq deconvolution, as well as 93 relevant immune-related genes. UPG PDGFRA-mutant GISTs appeared to have higher proportions of intratumoral CD4+ T cell and macrophage expression (Figure 3, middle), while UPG KIT-mutant GISTs primarily expressed CD4 and macrophage signatures only in the most immune cell-infiltrated tumors. Furthermore, B cell and CD8<sup>+</sup> T cells signatures were also expressed in GISTs with high ESTIMATE and CYT scores, confirming what we and others have shown regarding the immune cell prevalence in GIST (14, 27).

Differential gene expression (DGE) analysis revealed that 20 of 93 (21.5%) immune-related genes were significantly differentially regulated between UPG *KIT*- and UPG *PDGFRA*-mutant GISTs (bold indicates P < 0.05; Figure 3, bottom). Specifically, UPG *PDGFRA*-mutant GISTs expressed significantly more CCR5, BTLA, CD96, CD48, TNFRSF9, TNFSF8, CCR4, CXCL11,

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Figure 2. PDGFRA-mutant GIST is more immunologically active compared with KIT-mutant GIST. (A) ESTIMATE and CYT scores (left) and CD45 and CD8 normalized counts (right) of all KIT- and PDGFRA-mutant GISTs (n = 61; Supplemental Table 3). (B). ESTIMATE and CYT scores (left) and CD45 and CD8 normalized counts (right) of UPG KIT- and PDGFRA-mutant GISTs (n = 22; Supplemental Table 4). (**C**) GSEA showing multiple immune pathways enriched in UPG PDGFRA-mutant compared with UPG KIT-mutant GISTs. NES, normalized enrichment score. (D) ×10 magnification CD45 and CD8 IHC staining in KIT- and PDGFRA-mutant GISTs. Red text indicates specimen was included in RNA-Seq cohort, while samples represented in black text were not included. Representative samples of n = 6 per group are shown. (E) CD45 (top) and CD8 (middle) quantification of IHC staining. n = 6 per group. The number of CD45\* and CD8<sup>+</sup> cells per HPF was calculated by examining 5 HPFs per tumor and plotting the average per tumor. Bottom: CD45 expression by flow cytometry (for specimens in which flow cytometry data were available). \*P < 0.05, t test. Bars indicate median.

CXCR4, KDR, IL6R, TNFRSF8, TNFSF14, TIGIT, TNFRSF17, HLA-DQA2, CXCL14, and CXCL12. Meanwhile, *KIT*-mutant GISTs expressed significantly more TNFSF18 and MICA. Together, these data suggest that *PDGFRA*- and *KIT*-mutant GISTs elicit a different quality of immune infiltrate, which may ultimately provide driver-specific strategies for immunotherapy.

PDGFRA- and KIT-mutant GISTs have distinct signaling and cytokine signatures. To determine whether intracellular signaling or cytokine differences could be contributing to the observed differences in immune infiltration between UPG PDGFRA-mutant and UPG KIT-mutant GISTs, we utilized GSEA to detect differences in these pathways. On GSEA, we first observed enrichment of a known PDGFRA-mutant GIST signaling pathway gene set from a prior report analyzing 3 PDGFRA-mutant GISTs (Figure 4A, top) (30). Compared with UPG KIT-mutant GISTs, UPG PDGFRAmutant GISTs also demonstrated significant transcriptional activation of the PI3K signaling pathway (Figure 4A, bottom), which may have been driven by expression changes in PDGFRA, RELN, KDR, or ERK, 4 genes that have been found to be significantly differentially expressed between PDGFRA- and KIT-mutant GISTs not only in our cohort but also previously (30). Furthermore, many cytokine and chemokine pathways were significantly upregulated in UPG PDGFRA-mutant GISTs (Figure 4B), which we found to be related to increased expression of CXCL14, CCL7, CCL19, VEG-FA, and KITLG (Figure 4C).

The differences observed in *CXCL14* expression were particularly intriguing. First, *CXCL14* mRNA was expressed at significantly higher levels in nearly all UPG *PDGFRA*-mutant GISTs. Quantitative real-time PCR (qRT-PCR) of bulk human *PDG-FRA*- and *KIT*-mutant tumors also revealed that *PDGFRA*-mutant GISTs produced approximately 10 times more *CXCL14* mRNA than *KIT*-mutant tumors, and 21 times more *CXCL14* mRNA than the *KIT*-mutant GIST-T1 (Figure 4D, left) and GIST-882 cell lines (data not shown). Finally, *CXCL14* has been shown to contribute to NK<sup>+</sup>, CD4<sup>+</sup>, CD8<sup>+</sup>, and regulatory T cell chemotaxis and tumor regression in other tumor models (31, 32). UPG *PDGFRA*-mutant GISTs expressed *CXCL14* mRNA at levels higher than those shown to lead to immune-mediated tumor regression in HPV<sup>+</sup> head, neck, and cervical cancers, while *KIT*-mutant tumors expressed *CXCL14* mRNA at levels below this threshold (Figure 4D, right). Together,

these findings suggest that CXCL14 expression may contribute to the observed differences in immune infiltration between *PDG*-*FRA*- and *KIT*-mutant GISTs (31, 32).

PDGFRA mutation produces multiple HLA-diverse, strong binding neoepitopes. Given the association of neoantigen presentation with inflamed tumor/immune microenvironments in other malignancies, we hypothesized that PDGFRA-mutant GISTs may produce more potent neoepitopes when compared with *KIT*-mutant GISTs. To explore this, we first mapped out the 8-, 9-, and 10-mer neoepitopes produced by all driver mutations in our cohort, generating neoepitopes with the mutation placed at each amino acid position in the mutant peptide. Then, we tested the binding affinity of each neoepitope to the matched, patient-specific HLA type in which that mutation was found.

First, we wanted to determine whether total neoepitope burden and the number of high-affinity neoepitopes produced by each GIST specimen correlated with immune infiltration or cytolytic activity in GIST, since neoepitope burden has been shown to correlate with increased response to immunotherapy in a variety of cancers (33, 34). However, total neoepitope burden and the number of high-affinity (<500 nM binding) or very high-affinity (<50 nM binding) neoepitopes did not correlate with immune infiltration or cytolytic activity across all mutations in the GIST cohort (Figure 5A) or across *KIT*- and *PDGFRA*-mutant GISTs specifically (data not shown).

We observed that the majority of patients with an oncogenic mutation in PDGFRA or KIT produced at least one high-affinity neoepitope (<500 nM binding), regardless of mutation subtype (Figure 5B). We also found that the D842V mutation produced 34 unique, high-affinity neoepitope:HLA-specific binding combinations across the predicted HLA types in our cohort of 16 D842V-mutant patients (Figure 5B), while KIT exon 11 and other PDGFRA mutations on average produced only 2-3 per mutation (Supplemental Table 6). The D842V mutation generated 6 different neoepitopes that bound to 12 different HLA types in 14 patients in our cohort (Supplemental Table 7), one of which is the most common HLA type in the United States and present in 95.7% of White individuals (HLA\*A02:01) (35, 36), suggesting that this mutation could produce an immune response in a wide variety of patients, whereas KIT exon 11 neoepitopes bound to less-prevalent HLA types.

Since the number of patients with *D842V* mutations (n = 16) exceeded the number of patients with any individual KIT mutation (n = 1-4, Supplemental Table 7), and this may obscure our HLA observations, we explored mutation-specific neoepitope binding diversity more broadly against all HLA types validated by NetMHCPan3.0 (Figure 5C). Again, nearly all PDGFRA and KIT GIST mutations produced at least one high-affinity neoepitope capable of binding to many different HLA types. D842V neoepitopes bound with very high affinity (<50 nM binding) to HLA\*A02:01 and many other prevalent HLA types. Interestingly, 50% (4 of 8) of all PDGFRA mutations produced at least one very high-affinity neoepitope (<50 nM binding) to HLA\*A02:01, compared with only 14% of KIT mutations (2 of 14). Thus, while all KIT- and PDGFRA-mutant GISTs appear to produce highaffinity neoepitopes, PDGFRA-mutant GISTs appear to produce more high-affinity neoepitopes to more common HLA types.



**Figure 3. CIBERSORT and DGE analysis identify unique immune signatures in GIST.** CIBERSORT (middle) and immune gene expression (bottom) of UPG *KIT*- and UPG *PDGFRA*-mutant GISTs, organized by mutational driver and increasing ESTIMATE score (*n* = 22). Unsupervised row-normalized clustering of genes shows grouping of genes into distinct groups, suggestive of oncogene-driven immune profiles. Genes shown in blue were significantly enriched in UPG *PDGFRA*- compared with UPG *KIT*-mutant, while genes in green were significantly enriched in UPG *KIT*- compared with UPG *PDGFRA*-mutant samples. Enrichment was considered as an adjusted *P* < 0.1 as calculated by DESeq2 for R, while boldface for gene names indicates a significant difference, with an adjusted *P* < 0.05. Clinicopathologic characteristics of the UPG GIST specimens are shown in the annotation and in Supplemental Table 4.

Given that GIST is often driven by a single oncogenic mutation, and GIST harbors a rich immune infiltrate, this raises the possibility that oncogenic GIST mutations produce neoepitopes that may classify as what has recently been provocatively described as high-quality neoepitopes (37). Machine learning identifies immune gene signatures for GIST. Through bioinformatics prediction and biologic validation methods, it appears that in *PDGFRA*-mutant GIST, immune cells are more numerous and have higher cytolytic activity than in *KIT*mutant GISTs, and both subsets contain a unique immune pro-

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**Figure 4.** *PDGFRA-* and *KIT*-mutant GISTs have distinct signaling and cytokine signatures. GSEA showing enrichment of (**A**) *PDGFRA* signaling, PI3K signaling, and (**B**) cytokine signaling pathways in UPG *PDGFRA-* compared with UPG *KIT*-mutant GISTs (n = 22). (**C**) Distribution of cytokines between UPG *KIT-* and UPG *PDGFRA-*mutant GISTs, by RNA-Seq. Adjusted P < 0.05 as calculated by DESeq2. All data points are shown; boxes define the interquartile range, with whiskers extending to lowest and highest data points. (**D**) Left: Relative *CXCL14* mRNA expression by qRT-PCR in *KIT-* (n = 7) and *PDGFRA-*mutant (n = 7) GISTs from the RNA-Seq cohort, compared with the GIST-T1 cell line (expression set at 1; data not shown). Right: *CXCL14* mRNA expression relative to *GAPDH* × 10<sup>6</sup> in UPG *KIT-* and *PDGFRA-*mutant GISTs. Horizontal dotted line represents *CXCL14* mRNA expression needed to induce tumor regression (31). \*P < 0.05, *t* test. Bars indicate the median.

file. This makes characterizing a global immune infiltrate in GIST challenging and suggests that specific immunotherapy approaches may ultimately need to be targeted to the specific oncogenic mutation in GIST. We therefore sought to define the most important immune features differentiating *PDGFRA*- and *KIT*-mutant GISTs through machine learning, which is an unbiased approach not dependent on previously defined gene set analysis, allowing for novel and impartial observations. We specifically used all *KIT*-



**Figure 5.** *PDGFRA* **mutation produces multiple HLA-diverse, strong binding neoepitopes.** (**A**) Pearson's correlation of ESTIMATE (top) and CYT (bottom) scores with total neoepitope burden (left), number of high-affinity neoepitopes (middle), and number of very high-affinity neoepitopes (right) among all GIST samples (*n* = 75). (**B**) Left: Percentage of patients with the indicated mutation whose mutation produced a predicted high-affinity neoepitope. Right: Number of potential neoepitope:HLA binding events produced by mutation type, averaged over the number of mutations per group. (**C**) Heatmap of the binding affinities of *KIT* and *PDGFRA* mutation-specific neoepitopes to all validated NetMHCPan 3.0 HLA types. Clinicopathologic characteristics of the GIST specimens are shown in Supplemental Table 3. Additional details regarding mutations, neoepitopes, and HLA types used to create this heatmap are shown in Supplemental Tables 6 and 7.

and *PDGFRA*-mutant GISTs (Supplemental Table 3; n = 61), as well as only UPG *KIT*- and *PDGFRA*-mutant GISTs (Supplemental Table 4; n = 22) to develop 2 random forest models, and assumed that these immune differences would most likely be related to the oncogenic driver.

The combination of salient immune features capable of most accurately profiling a *PDGFRA-* or *KIT-*mutant GIST were identified based on each feature's "importance," which is calculated as part of caret for R's implementation of randomForest (38). The feature importance metric is a measure indicating how useful a feature is for separating samples into their distinct classes. The features with the highest importance provide the classifier with the highest increase in performance.

We included all 117 immune features to initially create the model (Supplemental Table 5) and subsequently retrained the model using fewer features. We noted that including more features in the model increased the model's performance on the training set, but decreased the model's performance on the test set, a phenomenon known as overfitting (Supplemental Figure 3A). Therefore, to prevent overfitting, we retrained the model using only the 6 most important features, which we believe will broaden the applicability of our model to future GIST samples. When we restricted the model to the 6 most relevant features (Figure 6A), 27 of 30 KIT-mutant GISTs were correctly classified as KIT-mutant in the training set, while 14 of 20 PDGFRA-mutant GISTs were correctly assigned as PDGFRA-mutant (sensitivity: 70%, specificity: 90%). The top immune features included CXCL14, TGF-BR1, TNFSF9, MICA, TNFRSF25, and CD96 (Figure 6B). Notably, the PDGFRA-mutant tumors that were incorrectly classified as KIT-mutant by our model had lower ESTIMATE and CYT scores than correctly classified PDGFRA-mutant tumors (Supplemental Figure 4A), while KIT-mutant tumors incorrectly classified as PDGFRA-mutant had more variable CXCL14 mRNA expression and higher TGFBR1 and CD96 mRNA expression than correctly classified KIT-mutant tumors (Supplemental Figure 4B). In our separate cohort of testing samples (n = 11), the 6-feature model correctly identified KIT- and PDGFRA-mutant GISTs with 91% accuracy (Figure 6C). To demonstrate the importance of these 6 immune features in classifying KIT- and PDGFRA-mutant GISTs, we excluded these features and retrained our random forest model, which reduced classifier performance and model accuracy from 91% to 72.7% (Supplemental Figure 3B).

We further validated our model on an external cohort of GISTs from the Complexity Index in Sarcomas (CINSARC) study (n = 12; ref. 39). Application of our immune profiling model to the CIN-SARC cohort correctly differentiated *KIT*- and *PDGFRA*-mutant GISTS with 83.3% accuracy (Figure 6C). Similar results were obtained when we created a model using only UPG *PDGFRA*-and *KIT*-mutant GISTs (Figure 6, D–F), in which CXCL14, IDO, TNFSF14, KDR, MICA, and TNFRSF9 were the most important features defining the model.

We employed the same machine learning approach to identify gene expression-based immune features associated with high PD-1 and PD-L1 expression across all GISTs (n = 75) in order to discover novel therapeutic targets and potential barriers to immune checkpoint blockade in GIST. Interestingly, our 6-feature model showed that increased CD27 expression was highly related to PD-1 upregulation (sensitivity: 90%, specificity: 87.1%, Figure 7A), along with PDCD1LG2, BTLA, CD40, CXCL14, and MICB, suggesting that these pathways may represent synergistic opportunities for PD-1 immunomodulation in GIST (Figure 7B). Using only these 6 features, our model was able to correctly predict high PD-1 expression in 92.9% of the testing cohort and 66.7% of the external CINSARC GIST cohort (Figure 7C).

PD-L1 mRNA expression correlated with PD-L1 protein expression in our cohort (Supplemental Figure 5; see complete unedited gel in the supplemental material). When training our PD-L1 random forest model, expression of key antigen-presenting machinery, including B2M, HLA.DRA, and TAP2, and the immune-responsive cytokines CXCL10 and LTA were predictive of increased PD-L1 expression across all GISTs (sensitivity: 76.7%, specificity: 87.1%, Figure 7D), suggesting a relationship between immune checkpoint ligand upregulation and immune recognition (Figure 7E). Applying the model of these antigen-presenting machinery features and immune cytokines, along with PD-L2, which was also identified as a feature predictive of high PD-L1 expression, our model correctly identified high PD-L1 expression with 85.7% accuracy in the testing cohort and 91.7% in the CIN-SARC GIST cohort (Figure 7F).

#### Discussion

The dramatic success of immune checkpoint blockade in a variety of notoriously difficult-to-treat cancers has nearly standardized immunomodulation as an approach for cancer treatment (40–42). However, subsequent research has advanced our understanding of how cancer type– and tumor cell–specific characteristics, such as genomic mutational burden, can predict response to immunotherapy (33, 34). It has also recently been shown that the state of the immune microenvironment can prevent response to immunotherapy if not appropriately addressed (43). In fact, it has become clear that the complex relationship between cancer cell–intrinsic factors and the subtleties of the cancer type–specific immune microenvironment may make a generalized strategy of immune checkpoint blockade ineffective (44). Therefore, it has become increasingly evident that cancer-specific immunotherapeutic strategies are needed.

In this study, we performed RNA-Seq on 75 human GIST specimens from 75 patients to characterize the immune landscape of different GIST mutational drivers, which we believe is the largest cohort of GISTs to be analyzed with next-generation RNA-Seq. After observing potential differences in immune infiltration, we focused on the 2 prevalent GIST mutational subtypes, namely *KIT* and *PDGFRA*, and found that *PDGFRA*-mutant GISTs contain more immune cells than *KIT*-mutant GISTs. This suggests that patients with *PDGFRA*-mutant GIST may ultimately have a greater potential to respond to immunotherapy.

Through differential gene expression analysis, we identified mutation-specific immune landscapes that may be significant for directing GIST immunotherapy. First, *PDGFRA*-mutant GISTs showed higher expression of CD96, suggesting that NK cells may play a more substantial role in the immune environment of *PDGFRA*-mutant compared with *KIT*-mutant GIST. In fact, NK cell infiltrate has been suggested to be a contributor to long-term survival in *KIT*-mutant GIST (45), which may help explain why *PDGFRA*-mutant GIST patients have a better over-



**Figure 6. Machine learning identifies an immune signature predictive of** *KIT***- and** *PDGFRA***-mutant GIST.** (**A**) Random forest modeling with 5-fold cross-validation of *KIT*- and *PDGFRA*-mutant GIST specimens (training set created by partitioning 80% of *KIT* and *PDGFRA* samples from Supplemental Table 3, *n* = 50). Confusion matrix (right) indicates assessment of model fit to training set. OOB, out-of-bag. (**B**) Distribution of top 6 features identified by random forest modeling. \*Adjusted *P* < 0.05 from DSeq2. (**C**) Predictive capacity of model on remaining *KIT*- and *PDGFRA*-mutant GIST testing set (*n* = 11) and the CINSARC cohort (*n* = 12). Accuracy (Acc), sensitivity, specificity, and *P* value[Acc >no information rate (NIR)] of the model are shown, calculated by caret package for R. Bars indicate mean ± SEM. (**D**) Random forest modeling with 5-fold cross-validation of UPG *KIT*- and UPG *PDGFRA*-mutant GIST specimens (training set created by partitioning 80% of UPG *KIT* and UPG *PDGFRA*-mutant GIST specimens (training set created by partitioning 80% of UPG *KIT* and UPG *PDGFRA*-mutant GIST specimens (training set created by partitioning 80% of UPG *KIT* and UPG *PDGFRA*-mutant GIST specimens (training set created by partitioning 80% of UPG *KIT* and UPG *PDGFRA*-mutant GIST specimens (training set. (**E**) Distribution of top 6 features identified by random forest modeling. \*Adjusted *P* < 0.05 from DSeq2. (**F**) Predictive capacity of model on remaining UPG *KIT*- and UPG *PDGFRA*-mutant GIST testing set (*n* = 4) and the CINSARC cohort (*n* = 12). Accuracy, sensitivity, specificity, and *P* value[Acc >NIR] of the model are shown, calculated by caret package for R. For **B** and **E**, all data points are shown, with boxes defining the interquartile range and whiskers extending to the lowest and highest data points.

all prognosis (4). Similarly, *PDGFRA*-mutant GISTs had higher CD4, CCR4, and CCR5 expression (indicative of regulatory T cells) and higher expression of the immunomodulators BTLA, CD48, TNFRSF9, and TIGIT. *PDGFRA*-mutant GISTs, there-

fore, may be more receptive to a variety of immunotherapeutic approaches, such as targeting TIGIT, BTLA, CD48, or TNFRSF9 in addition to regulatory T cells and NK cells, and PD-1/PD-L1 checkpoint blockade.

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**Figure 7. Machine learning identifies an immune signature predictive of PD-1 and PD-L1 expression in GIST.** (**A**) Random forest modeling of PD-1 with 5-fold cross-validation of GIST specimens (training set created by partitioning 80% of all GIST samples from Supplemental Table 1, n = 61). Confusion matrix (right) indicates assessment of model fit to training set. (**B**) Distribution of top 6 features identified by random forest modeling. \*Adjusted q < 0.1. (**C**) Predictive capacity of model on remaining 14 GISTs (testing set) and external CINSARC GIST cohort (n = 12). Accuracy, sensitivity, specificity, and *P* value[Acc > NIR] of model are shown, calculated by caret package for R. (**D**) Random forest modeling of PD-L1 with 5 k-folds cross-validation of GIST specimens (training set created by partitioning 80% of all GIST samples from Supplemental Table 1, n = 61). Confusion matrix (right) indicates assessment of modeling fit to training set created by partitioning 80% of all GIST samples from Supplemental Table 1, n = 61). Confusion matrix (right) indicates assessment of modeling fit to training set. (**E**) Distribution of top 6 features identified by random forest modeling. \*Adjusted q < 0.1. (**F**) Predictive capacity of model on remaining 14 GISTs (testing set) and external CINSARC GIST cohort (n = 12). Accuracy, sensitivity, specificity, and *P* value[Acc >NIR] of the model are shown, as calculated by caret package for R. For **B** and **E**, all data points are shown, with boxes defining the interquartile range and whiskers extending to the lowest and highest data points.

Comparison of UPG GISTs of the *KIT*- and *PDGFRA*-mutant phenotypes revealed that mitotic rate, an indicator of GIST aggressiveness, was inversely related to the quantity of the immune infiltrate. While this concept has been described in other cancers, tumor proliferative index has not yet been shown to affect the quantity of the immune infiltrate in GIST. *PDGFRA*-mutant GIST has been shown to have a lower proliferative index when compared with *KIT*-mutant GIST (46). However, when controlling for mutational driver, the relationship between mitotic rate and immune cell infiltration in UPG GISTs loses significance, suggesting that oncogenic driver may be more important for dictating the immune infiltrate. In fact, comparison of UPG *KIT*- and *PDGFRA*-mutant GISTs revealed distinct cytokine profiles, which we hypothesize may contribute to immune cell chemotaxis. CXCL14 is a relatively recently identified chemokine with a clear role in immune cell attraction (31, 32, 47). CXCL14 was found to be significantly upregulated in *PDGFRA*-mutant compared with *KIT*-mutant GISTs, not only in our cohort but also in a previously published cohort (30). Unfortunately, studying this correlation has proved challenging, as there is currently no *PDGFRA*-mutant GIST mouse model or human *PDGFRA*-mutant GIST cell line in which to explore what appears to be a clear biological relationship between PDGFRA tyrosine kinase activation and CXCL14 secretion. Thus, the implication of our finding at this time remains unclear.

It is clear that in cancers with a high number of mutations, tumor mutational burden correlates with immunotherapy response (33). However, the role of neoepitopes in the immune response of cancers with a single oncogenic mutation (i.e., GIST) has not been described. To our surprise, nearly every GIST mutation in our cohort produced a neoepitope capable of binding with high affinity to patient-specific HLA class I, suggesting that even cancers with one mutation can generate a neoepitope recognizable by the immune system. We discovered that the D842V mutation produced 6 different high-affinity neoepitopes, which bound to 12 different HLA types of patients in this cohort alone, one of which is the most common HLA type in the United States and in White individuals (35, 36). While we have not shown that these neoepitopes produced an immune response in these patients, it is intriguing that GISTs with a larger immune infiltrate appear to have a mutation that generates peptides with broad HLA binding specificity, suggesting that this mutation may be of higher quality (37). Moreover, the HLA-specific binding affinities of neoepitopes generated by mutation-specific GIST subtypes reported in this study may provide a road map for patient selection in future neoantigen vaccination trials.

We employed machine learning in order to identify key differences and to profile the expression of immune-related genes between PDGFRA- and KIT-mutant GISTs. Generating a gene expression-based immune profile that accurately predicted the KIT- or PDGFRA-mutant genotype not only supports our hypothesis that the genotype may impact the immune infiltrate, but also introduces another method by which profiling the immune landscapes of KIT- and PDGFRA-mutant tumors can lead to more personalized cancer immunotherapy. Not surprisingly, CXCL14 appeared to be a top feature of PDGFRA-mutant GIST, again suggesting that CXCL14 expression is oncogene specific and may be responsible for the observed differences in immune infiltration and activity. TGFBR1, TNFSF9, MICA, TNFRSF25, CD96, IDO, TNFSF14, KDR, and TNFRSF9 all contributed to the predictive capacity of the immune profiling models, which resulted in >90% diagnostic accuracy in our testing set and in 83% of the CINSARC validation set. We have previously shown the importance of IDO in dictating the immune phenotype of GIST (13). The role of NK cells and the additional immune modulatory receptors identified in this study should continue to be explored.

Since immunotherapy has not yet shown efficacy in advanced and metastatic GIST (48), we sought to explore the immune profiles of PD-1-high and PD-L1-high GISTs across our entire GIST cohort to identify both therapeutic opportunities and potential barriers to effective immunotherapy. CD27 was found to be an important feature predictive of high PD-1 expression in our random forest model, which correctly identified high PD-1 expression in 93% of our testing GIST cohort. Agonistic CD27 antibodies already exist and have shown efficacy with anti-PD-1 in other models, suggesting that this may be an effective strategy for GIST (49). Similarly, the costimulatory and checkpoint receptors CD40 and BTLA, along with the immune checkpoint ligand PDCD1LG2 and the MHC class I protein MICB were features predictive of high PD-1 expression. We have previously shown that CD40 ligation enhances antitumor immunity in our *Kit* exon 11 mouse model (17), and these other receptors should be considered in future trials.

Notably, applying our PD-1 model to the external CINSARC cohort resulted in only 67% accuracy, even though our PD-L1 model, in which the same methodology was used, was more accurate (92%). One reason may be related to the small sample size of the external cohort. However, these models identify high-priority targets for further experimental validation.

Finally, expression of key antigen-presenting machinery including B2M, PDCD1LG2, HLA-DRA, and TAP was found to be highly predictive of PD-L1 expression not only in our cohort (86% accuracy) but also in the external CINSARC GIST cohort (92% accuracy), suggesting a tumor cell-specific mechanism of linking neoepitope presentation with immune suppression. A similar finding has been observed in HIV-infected cells, where MHC class I machinery was upregulated concurrently with PD-L1 (50). This is an important concept to consider, as efforts to block PD-L1 may result in tumor cell changes that also downregulate MHC class I antigen presentation, which is essential for the CD8<sup>+</sup> T cell-mediated immune response. High PD-L1 expression has also recently been shown to independently correlate with a worse prognosis in soft tissue sarcomas, which was attributed to an increased degree of immune exhaustion, suppression, and negative regulation (51). In contrast, efforts to enhance antigen presentation using neoantigen peptide vaccination or GM-CSF-expressing tumor cell vaccines may require PD-L1 inhibition.

Ultimately, it is becoming increasingly evident that immunotherapy requires a specific approach targeted to not only the tumor type but also the tumor-specific immune microenvironment. Through RNA-Seq and 4 trained machine learning models, we have characterized important and consistent differences in the immune landscape of GISTs driven by different oncogenic mutations, which we believe may help guide future immunotherapy trials in GIST.

## Methods

*Human GIST specimens and clinicopathologic features.* Fresh surgical specimens were collected immediately upon resection and flash frozen in liquid nitrogen. RNA was extracted using the RNeasy kit (QIA-GEN) as per the kit-specified protocol. Clinicopathologic features were obtained via chart review of patient and pathologic records.

*Bioinformatics.* Poly(A) selection and next-generation RNA-Seq were performed on an Illumina HiSeq 2500 platform by the MSKCC Integrated Genomics Operations core facility. A minimum of 40–50 million 50-bp paired-end reads were obtained for each sample. RNA-Seq reads were processed using Trimmomatic v0.36 (52) using the following parameters recommended for paired-end Illumina sequenc-

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ing: PE - phred33, ILLUMINACLIP: TruSeq3-PE.fa:2:30:10, LEAD-ING: 3, TRAILING: 3, SLIDINGWINDOW: 4:15, and MINLEN: 36. Sequencing reads were then aligned to the human genome (version hg38\_r88), and gene-level counts were calculated using STAR version 2.6.0a (53). Read counts were then normalized, and DGE analyses were performed on indicated groups using the R software package DESeq2 (54, 55). GSEA was performed using the java GSEA software package (version 3.0) and Molecular Signatures Database (MSigDB) version 6.2 (Broad Institute) (56, 57).

The activity of cytotoxic T cells within each sample was estimated utilizing the CYT score, which was calculated from the geometric mean of the normalized read counts for perforin (PRF1) and granzyme A (GZMA) as previously described (23). The immune cell compartment of these samples was further quantified from the normalized read counts using the R package ESTIMATE (23). The ESTIMATE algorithm is composed of a nonimmune "stromal score" parameter and an "immune score" parameter. In this work, we refer to ESTI-MATE score as the immune score subcomponent of the ESTIMATE algorithm, which infers the degree of immune cell infiltration within tumor tissue based on expression data of 141 immune-related genes previously shown to correlate with the presence of leukocytes (24). The absolute quantities of individual immune cell subtypes was inferred from the normalized read counts by using the CIBERSORT program (29), which employs an externally validated leukocyte gene signature matrix (LM22) of 547 genes designed to distinguish 22 cell phenotypes including 7 T cell types, B cells, macrophages, monocytes, and NK cells (29). Heatmaps of gene expression data with corresponding clinicopathologic characteristics were created with the R package ComplexHeatmap (58). ssGSEAs were performed using the R software package GSVA (59).

Neoepitopes and HLA prediction. Patient HLA haplotypes were inferred using the program seq2HLA (60). For neoepitope identification, the amino acid sequences of patient-specific mutations were mapped to combinations of 8-, 9-, and 10-mer peptides, with the mutation evaluated at each amino acid position. Each patient-specific peptide was then tested with the respective patient-specific HLA type using NetMHCPan 3.0 to identify potential high-affinity neoepitopes (61, 62). Neoepitope burden was defined as any epitope with a predicted binding affinity within the top 2% when compared with 400 random natural peptides, as recommended by NetMHCPan 3.0. High-affinity neoepitopes were further filtered using a binding affinity threshold of 500 nM, while very high-affinity neoepitopes were defined as having <150 nM binding affinity. Finally, to explore mutation-specific neoepitope binding diversity, we tested mutation-specific neoepitopes more broadly against all validated NetMHCPan 3.0 HLA types and visualized them using heatmap.2 in R.

*Machine learning.* Random forest modeling with 5-fold cross-validation was performed on the training set using the software package caret for R to identify top predictors (38). One hundred and seventeen features, comprising normalized read counts of various immune genes, and bioinformatically inferred CIBERSORT scores were used to train the model (Supplemental Table 5). For characterization of *PDGFRA*-specific immune features, *KIT*- and *PDGFRA*-mutant GIST samples were first partitioned randomly into training (80% of samples) and testing sets (20% of samples). The training data set was used to train a random forest machine learning classifier using 5-fold cross-validation. During each of the 5 folds, a random forest model was trained on 80% of the training data, and training accuracy, sensitivity, and specificity were calculated; the remaining 20% of the training data were then used to calculate testing accuracy, sensitivity, and specificity. Finally, our trained machine learning model was applied to the 20% testing set and to the external validation CINSARC cohort (n=12; ref. 39) in order to assess model accuracy on a naive sample set.

Given the total sample size of our cohort and to prevent overfitting of our model, random forest modeling with 5-fold cross-validation was re-performed on the training set using only the top 6 identified features, and the model was then applied to the naive testing set and CINSARC external data set to assess for model accuracy. Feature importance is a variable calculated by the caret package, with a higher number indicating that the feature is more important for model accuracy (38). The confusion matrices shown in Figures 6 and 7 reflect the model, with the highest accuracy after 5-fold cross-validation and optimization by caret. The P value used by caret is calculated using an exact binomial test in which a 1-sided test is performed to determine whether the model accuracy rate is better than the "no information rate," which represents the proportion of data within the majority class. For generation of PD-1- and PD-L1-specific immune features, the same approach was used, with groups split into high and low expression based on median PD-1 (median: 14.1) or PD-L1 (median: 209.5) normalized read counts in the training set.

*qRT-PCR*. Total RNA was extracted from snap-frozen human tumors, reverse transcribed, and amplified with PCR TaqMan probe for *CXCL14* (Hs01557413\_m1). qRT-PCR was performed using a ViiA 7 real-time PCR system (Applied Biosystems). Data were calculated by the  $2^{-\Delta\Delta Ct}$  method as described in the manufacturer's protocol and were expressed as fold increase over the GIST-T1 or GIST-882 cell line control (63, 64).

*Flow cytometry*. Flow cytometric analysis was immediately performed at the time of specimen collection on freshly obtained human GIST specimens after tumor dissociation as previously described (15). Cells were analyzed on a BD FACSAria or LSRFortessa (BD Biosciences). A human-specific antibody for CD45 (2D1) was obtained from BD Biosciences.

Western blot analysis. Protein from flash-frozen GIST tissue or cell lines was analyzed as described previously (18). Antibodies used against GAPDH (clone D16H11) and PD-L1 (clone E1L3N) were purchased from Cell Signaling Technology.

*Histology.* Freshly collected tumors were fixed in 4% paraformaldehyde, embedded in paraffin, and cut into 5- $\mu$ m sections. CD45 (PD7/26 + 2B11) and CD8 (SP57) staining was performed by the MSK-CC Molecular Cytology Core. Slides were scanned with MIRAX SCAN (Zeiss) and photographed with Pannoramic Viewer (3DHISTECH Ltd.). For quantification of CD45 and CD8 staining, cell counting was performed manually in a blinded fashion at ×20 magnification (HPF). The number of CD45<sup>+</sup> and CD8<sup>+</sup> cells per HPF was calculated by examining 5 HPFs per tumor and plotting the average per tumor.

*Statistics*. Some data and graphs were created using Prism 7.0 (Figure 2, A, B, and E; Figure 4, C and D; Figure 5, A and B; Figure 6, B and E; Figure 7, B and E; GraphPad Software). Statistical tests were performed as described in the figure legends, and when possible, adjusted *P* values calculated by DESeq2 were used to assess for significance, with a threshold of adjusted P < 0.05. For Figure 7, B and E, an FDR approach was taken using the Benjamini, Krieger, and Yekutieli 2-stage step-up method, with a *q* value threshold of <0.1 in order to assess for significance.

Data availability and CINSARC external validation GIST cohort. RNA-Seq data presented in this work are available through the Sequencing Read Archive under accession no. PRJNA521803. RNA-Seq data of fresh-frozen GIST tumors from the publicly available CINSARC cohort (39) were obtained from the Sequence Read Archive under accession no. SRP057793.

*Study approval.* Tumor specimens were obtained from 2011 to 2017 from 75 GIST patients who underwent surgical resection at MSKCC. Prior to resection, patients provided written informed consent under an IRB-approved protocol. Using a prospectively maintained institutional database, clinicopathologic characteristics were documented and encrypted, and all research was conducted in compliance with Health Insurance Portability and Accountability Act regulations.

## Author contributions

GAV, TGB, BDM, and RPD contributed to study conception and design. GAV, TGB, and RPD contributed to the development of research methodology. GAV, TGB, BDM, JQZ, JKL, NJP, SZ, ML, and FC contributed to the acquisition of data and conduction of

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experiments. GAV, TGB, ML, RLG, NJP, FR, and RPD contributed to the analysis and interpretation of data. GAV, TGB, and RPD contributed to the writing, review, and revision of the manuscript. All authors reviewed and contributed comments that were incorporated into the final version of this manuscript. Study supervision was provided by RPD.

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