Development of a cross-platform biomarker signature to detect renal transplant tolerance in humans

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Identifying transplant recipients in whom immunological tolerance is established or is developing would allow an individually tailored approach to their posttransplantation management. In this study, we aimed to develop reliable and reproducible in vitro assays capable of detecting tolerance in renal transplant recipients. Several biomarkers and bioassays were screened on a training set that included 11 operationally tolerant renal transplant recipients, recipient groups following different immunosuppressive regimes, recipients undergoing chronic rejection, and healthy controls. Highly predictive assays were repeated on an independent test set that included 24 tolerant renal transplant recipients. Tolerant patients displayed an expansion of peripheral blood B and NK lymphocytes, fewer activated CD4+ T cells, a lack of donor-specific antibodies, donor-specific hyporesponsiveness of CD4+ T cells, and a high ratio of forkhead box P3 to α-1,2-mannosidase gene expression. Microarray analysis further revealed in tolerant recipients a bias toward differential expression of B cell–related genes and their associated molecular pathways. By combining these indices of tolerance as a cross-platform biomarker signature, we were able to identify tolerant recipients in both the training set and the test set. This study provides an immunological profile of the tolerant state that, with further validation, should inform and shape drug-weaning protocols in renal transplant recipients.

Introduction

Transplantation tolerance can be defined as the stable maintenance of good allograft function in the sustained absence of immunosuppressive therapy. In the clinical arena, it is only apparent when patients experience stable allograft function despite having ceased all immunosuppression for an extended period of time. This state, defined as operational tolerance, has rarely been reported in renal transplantation (1–5), being more apparent when patients experience stable allograft function despite having ceased all immunosuppressive therapy. In the clinical arena, it is only

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Results

Tolerant renal transplant patient demographics. The training set comprised 71 European kidney transplant recipients and 19 age- and sex-matched healthy controls (Table 1). The Tol-DF group had a high percentage of cadaveric donors (7 of 11), a high degree of HLA mismatching (median mismatches, 4.0), was predominantly male (9 of 11), and had varied causes of end-stage renal failure and some evidence of sensitizing events, such as blood transfusions and previous transplants (Table 2). These patients had relatively uneventful post-transplantation courses, with only 1 patient having a documented episode of acute cellular rejection (ACR). The period of being immunosuppression-free varied from 1 to 21 years.

The Tol-DF group of the test set (Table 3) consisted of 24 patients, most of whom had received their transplant from a highly HLA-matched living donor (median mismatches, 0.0) and had ceased taking all immunosuppression medication for periods from 1 to 32 years.

Tol-DF recipients displayed increased numbers of B and NK lymphocytes. As shown in Figure 1, Tol-DF patients of the training set displayed an increased percentages of peripheral blood B and NK cells and a corresponding decrease in the percentage of T cells. When the percentages of B cells and T cells were expressed as a ratio, Tol-DF patients displayed the highest ratio compared with all other study groups, including HCs. For 6 Tol-DF patients and 10 s-CNI patients, it was also possible to calculate the absolute number of cells per lymphocyte subset. This showed that the altered ratio was
due to an expansion in B and NK cell numbers and not a reduction in T cell numbers, as none of the Tol-DF group were lymphopenic (Table 2).

In line with our findings in the training set, Tol-DF patients of the test set also showed elevated percentages of peripheral blood B cells and a higher ratio of B/T cell percentages (Figures 1, E and H) compared with all other groups except HCs.

Given the distinct increase in peripheral blood B cells detected in Tol-DF patients, B cell subsets were analyzed (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI39922DS1) and cytokine production (Supplemental Figure 2) assessed in selected patients of both study sets. The Tol-DF group displayed a trend toward redistribution of B cell subsets, with a decreased memory pool and concomitant increase in transitional and naive B cell subsets. When the percentages of B cell subsets were examined as a ratio, Tol-DF patients were found to have a significantly lower proportion of memory and higher proportion of transitional B cells compared with CR patients. A significant proportion of B cells from Tol-DF patients was found to produce TGF-β upon in vitro stimulation, rather than IL-10 or IFN-γ. However, no significant differences in production of IL-10 were detected for any study group. The capacity of B cells from each patient group to produce either cytokine on stimulation was analyzed by calculating the ratio of the number of B cells producing each cytokine. This suggested that B cells of Tol-DF patients had a skewed cytokine response, with a higher propensity for TGF-β production than B cells from other study groups.

**Tolerant recipients had fewer activated CD4+ T cells in peripheral blood.** Expression of CD25 by CD4+ T cells was analyzed as described in Methods. Tol-DF patients in the training set were found to have significantly lower percentages of circulating CD4+CD25hi T cells, broadly thought of as activated T cells (10, 11) (Figure 2A), compared with the HC, s-LP, s-nCNI, and CR groups. Interestingly, no significant differences in the percentages of CD4+CD25hi Tregs were detected among the study groups (Figure 2B). Similar results were also found in the test set, with Tol-DF patients having significantly lower percentages of CD4+CD25hi T cells compared with s-CNI and chronic allograft nephropathy (CAN) groups, but again no differences in the percentages of CD4+CD25hi Tregs were detected between Tol-DF and any other study group (Figure 2, C and D). Statistical comparisons between other groups of the training and test sets are shown in Supplemental Table 1, A and B, respectively.

When we tested the ability of enriched CD4+CD25hi T cells to suppress autologous T cell proliferation induced by polyclonal stimulation, no significant differences were found between any of the patient groups or HCs (data not shown). Furthermore, Tol-DF patients did not display higher percentages of...
The majority of tolerant recipients did not have detectable anti-donor HLA-specific antibodies. Serum non-donor-specific antibodies (NDSAs) were detectable in some patients from all study groups of the training set (Figure 3A) by Luminex xMAP analysis. Within this cohort, no Tol-DF patients had detectable donor-specific antibodies (DSAs), whereas all other groups had some patients with detectable DSAs, with almost half of the CR patients having detectable levels of both donor- and non-donor-specific anti--HLA class I and class II antibodies. Similar to the training set, only 1 of 22 Tol-DF patients within the test set had detectable DSAs (data not shown).

Interestingly, in general, graft function was worse in DSA-positive patients than DSA-negative patients, with an estimated glomerular filtration rate (eGFR) of 31 (range, 17–87) in DSA-positive patients compared with 60 (range, 13–94) in DSA-negative patients. The possible pathogenicity of detected anti-donor antibodies was tested in the training set (Figure 3B). In 7 of 20 patients with anti-class I antibodies and 4 of 13 patients with anti-class II antibodies, we found complement-fixing isotypes (IgG1 and IgG3); the remaining positive cases were exclusively of non-complement-fixing isotypes.

Detection of non-donor-specific anti-class I and anti-class II antibodies was significantly associated with having received a previous transplant and having detectable panel reactive antibodies before transplant (Fisher’s exact test, P < 0.05), but not with previous pregnancies, blood transfusions, graft dysfunction, or episodes of ACR. In contrast, donor-specific anti-class I antibodies were associated with previous episodes of ACR and the number of HLA mismatches between donor and recipient (Fisher’s exact test, P < 0.05).

Tolerant patients had lower frequencies of direct pathway anti-donor IFN-γ CD4⁺ T cell responses. Comparison of direct pathway CD4⁺ T cell anti-donor and anti-third party (equally mismatched to donor) responses was assessed by IFN-γ ELISpot. Tol-DF patients had significantly higher ratios of responder anti-donor/anti-third party frequencies, indicating donor-specific hyporesponsiveness, compared with all other stable patient groups within the training set (Figure 4A; individual responder frequencies against donor and third party are shown in Supplemental Figure 3). Donor-specific hyporesponsiveness was not mediated by Tregs, as depletion of CD25⁺ cells from responder T cells did not result in an increase in responder frequencies (data not shown).

As patients within the Tol-DF group of the test set were frequently completely HLA matched with their donors, anti-donor and anti-third party IFN-γ responses were generally very low (responder frequencies, >1,200,000). Despite this, the trend in anti-donor responses in this Tol-DF group was generally reproduced, although a significant difference compared with other groups was not detected (Figure 4B).

Tolerant recipients displayed a higher ratio of expression of FoxP3 to α-1,2-mannosidase genes in peripheral blood. Whole blood gene expression levels of FoxP3 and α-1,2-mannosidase, both of which have been shown to correlate with anti-donor immune reactivity after transplantation (12), were analyzed by quantitative RT-PCR (qRT-PCR) (Supplemental Figure 4). When calculating the ratio of FoxP3 to α-1,2-mannosidase expression, a significant difference was detected between Tol-DF and the CR and HC groups of the training set (Figure 5A). The patient groups displaying the highest ratio were HC, s-LP, and Tol-DF, whereas the ratio was dramatically lower in CR patients (Mann-Whitney U test P values for comparisons between groups other than Tol-DF of the training and test sets are shown in Supplemental Table 2, A and B, respectively). This ratio significantly correlated with eGFR and inversely correlated with serum CRT (Pearson coefficients: 0.372, P = 0.002, and -0.299, P = 0.014, respectively; data not shown).

When the same analysis was performed on the test set, the ratio in Tol-DF patients was significantly higher than in all other patient groups except HCs (Figure 5B). Combining the training and test set observations shows that tolerance is associated with a high ratio of peripheral blood FoxP3 to α-1,2-mannosidase expression.

Tolerant patients exhibited a distinct gene expression profile. The RISET 2.0 custom microarray, designed with a focus on transplantation research, was assembled by the inclusion of 5,069 probes and used to analyze the expression of 4,607 genes (valid Entrez Gene ID) in peripheral blood samples.

A 4-class analysis of microarray data was performed on the training set (Figure 6). Significantly altered gene expression detected between Tol-DF patients and other comparator groups, stable recipients (s-CNI, s-nCNI, and s-LP), CR patients, and HCs, was statistically determined using the Kruskal-Wallis non-parametric test with adjustment for false discovery rate (FDR) at 1% (13). The HC group was included in this analysis in order to address the lack of immune suppression in Tol-DF patients compared with the other study groups. Two hundred and sixty probes, corresponding to 255 genes, were identified as being significantly differentially expressed between the study groups (Supplemental Table 3A; median fold changes in gene expression for each patient group are shown in Supplemental Figure 5A). When a similar analysis was performed on the test set, 1,378 probes, corresponding to 1,352 genes, with significantly altered expression were identified (Supplemental Table 3B and Supple-

### Table 3

<table>
<thead>
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<th></th>
<th>Age³</th>
<th>% Female⁸</th>
<th>Post-Tx⁶</th>
<th>CRT⁰</th>
<th>wbc⁵</th>
<th>% 1st Tx⁷</th>
<th>HLA-MM⁹</th>
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<td>51</td>
<td>(45–60)</td>
<td>36</td>
<td>19</td>
<td>13–29</td>
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<td>36</td>
<td>12</td>
<td>0–38</td>
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<td>44</td>
<td>(39–55)</td>
<td>45</td>
<td>6</td>
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<td>–</td>
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<td>40</td>
<td>8</td>
<td>0–42</td>
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Median and interquartile ranges are shown. ³Age in years. ⁸Percentage of females. ⁶Time after transplantation (yr). ⁰Serum CRT values (normal range, 80–105 μmol/l); ⁵wbc count (×10⁹ cells/l). ⁷Percentage of patients with their first transplant. ⁹Median number of HLA-A, -B, and -DR mismatches between donor and recipient (maximum, 6).
Differential gene expression detected by microarray analysis was reproducible by qRT-PCR. Gene expression detected by microarray was reproducible by qRT-PCR.

Gene expression diagnostic capabilities. For a more precise quantitative approach to gene expression analysis, with the utility to distinguish tolerant from non-tolerant individuals, we employed the top-ranked genes identified by microarray analysis, excluding any overlapping probes for any single gene (e.g., TCLA-1 ranked 2 and 4, excluding probe ranked 4), in an additive binary regression model to build receiver operating characteristic (ROC) curves. These probes were used to build a gene expression signature to specifically identify Tol-DF patients by first producing predicted classes (within-sample) and hence a classification for each individual.

For this analysis, 2-class ROC curves (tolerant versus non-tolerant) were built by both including and excluding the HC group from the nontolerant comparator groups. This was done because while the comparison of HCs with tolerant individuals is of interest in identifying tolerance-specific gene expression, in the context of developing a clinical diagnostic test for tolerance in renal transplant patients, this comparison is not useful.

The corresponding ROC curve built excluding HCs (Figure 7A) and based on the expression of the top 10 ranked genes (Table 4) delivered a peak specificity and sensitivity of 1, with a threshold of 0.01, and corresponding positive predictive and negative predictive values (PPV and NPV, respectively) of 100% within the training set (ROC including HC: threshold 0.2, specificity 0.853, sensitivity 0.923). Although 6 genes were sufficient to deliver good discrimination of tolerant patients within the training set, the top 10 ranked genes were selected for use, as they improved the specificity and sensitivity of subsequent ROC analysis of the test set (Figure 7B). Within sample analysis of the test set delivered a peak specificity of 0.890 and sensitivity of 0.806, with a threshold of 0.35, and PPV and NPV of 71% and 93%, respectively (ROC including HCs: threshold 0.23, specificity 0.801, sensitivity 0.806).

Annotation enrichment analyses on significant genes coincide between the training set and the test set. To identify significant associations of tolerance-related genes with any specific molecular pathway screened by microarray, we performed annotation enrichment analyses on the set of 174 overlapping probes identified between the training and test sets. The majority of genes found to have any significant association with annotated pathways were enriched within B cell–related pathways (Supplemental Table 6). In line with these data, of the top 11 ranked probes, corresponding to 10
genes, 6 genes are described to be expressed by B cells or related to B cell function (Table 4). This generalized B cell signature of tolerance is also cross-validated by Newell et al. (9). In addition to the B cell–related pathways enriched within this probe list, other pathways were also significantly regulated, including those involving protein tyrosine kinases and generation of secondary signaling messenger molecules as well as other T cell activation–related pathways (Supplemental Table 6).

Cross-platform biomarker diagnostic capabilities. All assays described in Methods section were tested in parallel for their diagnostic ability to distinguish Tol-DF patients from all other study groups. Assays performed on the test set were those that were highly predictive of tolerance within the training set and are shown above. By combining bioassays and biomarkers that indicate the presence of tolerance, we would expect that by using a cross-platform approach, we can significantly improve the diagnostic ability of any such individual test. This was indeed observed for the test set.

Indeed, when biomarkers and microarray data were analyzed in combination, Using (a) the ratio of B to T lymphocyte subsets, (b) the percentage of CD4+CD25int T cells, (c) the ratio of anti-donor to anti–third party ELISpot frequencies, (d) the ratio of FoxP3 to α-1,2-mannosidase expression, and (e) a signature of the top 10 ranked genes, the specificity and sensitivity for the training set was 1, with a threshold of 0.01, which implied PPV and NPV of 100% (Figure 8A).

When analyzing the test set, a peak specificity of 0.923 and a sensitivity of 0.903 were obtained, with a threshold of 0.27, and a corresponding PPV of 80%, and NPV of 96% (Figure 8B), which improved the diagnostic capacity compared with that obtained with gene expression alone.

Therefore, application of a cross-platform biomarker signature may be more capable of identifying bona fide tolerance, as in addition to gene expression and phenotype analysis, it can also take into consideration an individual’s immunological functional state, which may be more closely related to describing the mechanistic basis of tolerance.
The utility of this cross-platform biomarker signature lies in its ability to identify renal transplant patients who may be unknowingly operationally tolerant. As shown in Figure 8, C and D, 5 stable recipients of the test set could be identified to have the tolerant signature and therefore may benefit from managed weaning from immunosuppression. Interestingly, 2 CAN patients of the test set were also identified as having a high probability of being tolerant. This finding may be explained by differences in the clinical assessment of chronic rejection, as unlike the CR group of the training set, CAN patients were not proven by biopsy to have immunemediated rejection, but were defined on the basis of poor graft function. It is possible that the cross-platform biomarkers used to test these patients have sufficient sensitivity to detect subtle differences between these 2 patient groups, a property that may be revealed by serial immune monitoring of patients such as these over time.

Discussion
In this study we have developed a cross-platform set of biomarkers that distinguish tolerant renal transplant recipients from patients with stable renal function under different degrees of immunosuppression, patients undergoing chronic rejection, and HCs. Biomarkers identified in a training set of tolerant patients have been validated in an independent test set.

Both this study and that of Newell et al. (9) have found an expansion of B and NK cells in peripheral blood of drug-free tolerant recipients, which is similar to the findings of a previous study on a smaller cohort of similar patients (14). Not only do our findings extend earlier observations, but they also highlight the value of an altered T/B cell ratio as a biomarker of transplantation tolerance. Microarray analysis also revealed a clear and strong B cell bias of genes with altered expression between Tol-DF and the other groups. Whereas the role of T cells in initiating and maintaining allograft rejection (15, 16) and tolerance (17) has been clearly established, the role of B cells and the mechanisms whereby they may contribute to the tolerant state have yet to be elucidated. Interestingly, a murine study of transplantation tolerance induced by anti-CD45RB therapy has shown a mechanistic role for B cells (18). Recent data have also shown the ability of naive B cells, following antigen-specific cognate interactions, to induce Tregs that inhibit graft rejection in a murine model of heart transplantation (19). While no significant increase in Br-1 (IL-10–producing B cells) was detected in any patient group within this study, the data presented here show altered ratios of B cell transitional and memory populations, a relative increase in TGF-β–producing B cells, and absence of serum donorn-specific antibodies and donor-specific direct T cell hyporesponsiveness in tolerant recipients. These observations allow speculation that renal transplant tolerance may be associated with alterations in both T cell- and B cell–mediated functions. Given the essential role of antigen-specific B cell–T cell help, this would be an anticipated finding, although no correlation between anti-donor IFN-γ ELISpot responder frequencies and the presence/absence of donor-specific antibodies was detected within other study groups (data not shown). However, Porcheray et al., studying both B cell and T cell immunity in combined kidney and bone marrow transplant recipients, recently demonstrated the uncoupling of T cell and B cell anti-donor immunity in some of their tolerant patients.

Figure 4
Detection of donor-specific hyporesponsiveness in renal transplant study groups by IFN-γ ELISpot analysis. IFN-γ ELISpots were used to detect direct pathway alloresponses in patients of the (A) training and (B) test sets. The number of IFN-γ–producing cells in recipient CD4+ T cells was calculated (background-deducted) when stimulated with donor cells and third-party cells (3rdP), to obtain a frequency of responder cells (1 responder/n cells). Box plots show median and interquartile range. Whiskers above and below the boxes indicate the 5th and 95th percentiles. Ratio of responder frequencies on donor/3rdP stimulation are shown. Ratio values greater than 1.5 indicate hyporesponsiveness to donor. Two-sided P values for Mann-Whitney U test comparisons between groups are shown (**P < 0.01, *P < 0.05). Individual patient IFN-γ ELISpot responder frequencies to donor and third party are shown in Supplemental Figure 3. γ, Wilcoxon test between donor and 3rdP frequencies, P < 0.05.

Figure 5
Ratio of FOXP3 to α-1,2-mannosidase (MAN1A2) expression assessed by qRT-PCR. A ratio of the expression values of FOXP3 and MAN1A2 in peripheral blood, determined by qRT-PCR, was calculated for patients of the training set (A) and test set (B). Box plots show median and interquartile range. Whiskers above and below the boxes indicate the 5th and 95th percentiles. Two-sided P values for Mann-Whitney U test comparisons between Tol-DF and other groups are shown (**P < 0.001, *P < 0.01). Statistical values for comparisons between other study groups are shown in Supplemental Table 2, A and B.
with tolerant renal transplant patients. An additional 8 genes with highly differential expression, overlapping between the training and test sets, were common to several B cell–related genes identified by Newell et al., by their comparison of tolerant patients with stable or HC groups (FCRLA, IGHM, IGVK3, EBI2, CD40, BLNK, CD79A, CD79B) (9). The detection of significantly altered B cell–related gene expression by this study is further reinforced by the enrichment of genes with significantly different expression being found to predominantly associate with B cell–related pathway annotations. Although both this and the Newell study show a clear B cell bias in differentially expressed genes, the limited overlap between the most predictive genes of tolerance identified by our studies may be attributed to the distinct methods of RNA sample preparation and data analysis used (28).

A possible interpretation of the tolerant signature described by this study could be that the immunological biomarkers detected are merely due to the lack of drug-mediated immune suppression in the Tol-DF group. To address this possibility, the study groups of the training set were specifically selected to include stable renal transplant patients on distinct immunosuppressive regimes and HCs as immune suppression-free subjects. Although clear differences between the HC and Tol-DF groups were observed in the training set, these differences were not reproduced in the test set, a finding that may be attributed to the fact that the mechanisms of tolerance may be more subtle within the test set, where tolerant recipients are highly HLA matched to their donors, in contrast to the training set.

As all of these study groups have been taken into consideration, the cross-platform signature described here appears to be a specific description of transplant tolerance, rather than simply a consequence of the absence of immunosuppression. It is pertinent to observe that while detailed comparison of tolerant patients and HCs may reveal more about the mechanistic basis of tolerance, in the clinical context, this comparison is not entirely relevant.

An interesting comparison is that of Tol-DF and s-LP patient groups of the training set, which differ in the use of 10 mg/d prednisone, considered by many clinicians as quasi-physiological. The s-LP group had a higher proportion of female recipients, a higher percentage of cadaveric donors, and poorer kidney function than the Tol-DF group. Rather counterintuitively, in most of the assays described, there are clear differences between these 2 groups in immunophenotype, anti-donor responses, FoxP3/α-1,2-mannosidase ratio, and gene expression. This supports the notion that steroid monotherapy can induce a significant difference in the patient’s immune status that can be evidenced by biomarkers.

One of the Tol-DF patients within the training set had received a bone marrow donation 4 years prior to kidney transplantation from the same donor. Immune suppression was initially withdrawn from this patient, as evidence of chimerism was detected. As the mechanisms of tolerance induction could be different in this patient, biomarker and ROC curve analysis were performed with inclusion and exclusion of this patient; however, this patient

(20). In this respect, the B cell signature observed in tolerant renal patients in this study may indicate an important role for B cells in promoting tolerance, or it may simply be an epiphenomenon. Nonetheless, this and the study by Newell et al. (9) indicate that more research focus on B cells in transplantation is warranted.

Monitoring of anti-donor responses using functional assays has demonstrated that hyporesponsiveness of direct pathway T cells develops over time after solid organ transplantation (21, 22). In the clinical context, enumerating the frequency of anti-donor T cells has proven useful in steroid withdrawal protocols (23).

In our study, measuring anti-donor direct pathway responses by ELISpot has also proven useful, where determining the ratio of responses against donor and third party reveals donor-specific hyporesponsiveness in tolerant patients. This test, however, is more useful when donor and recipient have several HLA mismatches.

Similar studies to this have focused on gene profiling of tolerant liver (24, 25) and also tolerant kidney recipients (26, 27). The set of genes that were differentially expressed in those studies and in the study reported here have little overlap. This possibly reflects differences in the organ, the patient groups, the RNA source and preparation protocol, or the analysis platform used. Indeed, the microarray used in this study was selectively designed based on both published and unpublished data to have a transplantation focus and therefore included a substantial number of immune response–related probes.

Of the genes identified by Martínez-Llordella et al. as being associated with the gene profile of tolerant liver transplant recipients (24), 2 genes, KLRF1 (NKp80), expressed on all NK cells, and CLIC3 (chloride intracellular channel 3), were found to be highly differentially expressed within the training and test sets of this study. Interestingly however, 2 of the most highly ranked genes associated with tolerance found within the training set, TCL1A (rank 2) and MS4A1 (CD20) (rank 5), are both B cell–related genes and have also been described by Newell et al. (9). Furthermore, MS4A1 has previously been identified by Braud et al. (26) as being associated with stable or HC groups (9). The top 10 ranked probes that overlapped with genes identified in the test set were subsequently used for ROC analysis.

Figure 6

Algorithm for microarray gene expression analysis. Four-class analysis of microarray gene expression data identified probes significantly differentially expressed between all patient groups of the training and test sets using a Kruskal-Wallis nonparametric test. Probes were ranked within the training set based on their P values with adjustment for 1% FDR. The top 10 ranked probes that overlapped with genes identified in the test set were subsequently used for ROC analysis.
did not appear as an outlier within the tolerant group in any of the assays in the study.

The utility of this tolerant signature depends on its ability to identify transplant recipients that can safely be weaned from immunosuppression. We have now developed a specific and sensitive set of reproducible assays that, when combined, can identify tolerant renal allograft recipients and also several renal transplant recipients on immunosuppressive drugs. Validation of these biomarkers and bioassays has been achieved using a completely independent set of patients, and this validation is reinforced by the fact that the test set was derived from a genetically different population and that there were also differences in the collection and processing of test set and training set samples. However, before these cross-platform biomarkers can be implemented as a decisional tool in the clinical setting, the findings of this study are being carried forward into a larger observational clinical study of immune monitoring in an independent cohort of renal transplant recipients (www.transplant-tolerance.org.uk). This signature provides a model for future immunosuppression minimization protocols, which in combination with immune monitoring of emerging biomarkers of rejection (29, 30) may allow for tailored and safe clinical posttransplantation management of renal allograft recipients.

**Methods**

*Training set description.* This cohort of patients, recruited by the IOT network, consisted of 71 kidney transplant recipients and 19 age- and sex-matched HCs. Five patient groups were included: 11 functionally stable kidney transplant recipients (serum CRT, <160 μmol/l; and <10% rise in the last 12 months) despite having stopped all their immunosuppression for more than 1 year (Tol-DF); 11 patients with stable renal function (same criteria) maintained on less than 10 mg/d prednisone as the only immunosuppressive agent (s-LP); 10 patients maintained on “full” immunosuppression (azathioprine and prednisone) in the absence of a calcineurin inhibitor since transplantation (s-nCNI); 30 patients maintained on standard calcineurin inhibitor therapy (s-CNI); 9 patients with biopsy-proven (all reevaluated for this study) and immunologically driven chronic rejection (CR). Patient clinical characteristics are described in Table 1. eGFR was calculated from serum creatinine by the Modification of Diet in Renal Disease (MDRD) equation (http://nephron.org/mdrd_gfr_si).

Approval of the study protocol was obtained from the Hammersmith and St Mary’s Research Ethics Committee (REC), which was the main study REC (application: 2002/6378), and from the RECs of all other European recruitment sites. All participants in this study provided informed consent. All samples were processed and analyzed in a blinded fashion.

*Test set description.* An independent set of kidney transplant recipients were recruited in the United States through the ITN study; some of these patients were included and described by Newell et al. (9). The protocol was approved by the institutional review board of each participating center and by a data safety monitoring board convened by the NIAID. The cohort consisted of (a) Tol-DF (n = 24) patients, functionally stable kidney transplant recipients (serum CRT within 25% of baseline) despite having stopped all immunosuppression for more than 1 year; (b) “Mono” patients (n = 11) with stable renal function who were maintained on monotherapy with steroids; (c) s-CNI subjects (n = 34), with clinically stable renal function using the same criteria as Tol-DF patients while on maintenance with a triple drug immunosuppressive regimen (including a calcineurin or mTOR inhibitor, an anti-proliferative agent, and corticosteroids); and (d) CAN participants (n = 20), defined as those with chronic allograft nephropathy
### Table 4
List of top-ranked significant genes within the training set and their annotation enrichment

<table>
<thead>
<tr>
<th>Gene rank</th>
<th>Gene</th>
<th>Gene symbol</th>
<th>Official symbol</th>
<th>Description</th>
<th>Relevant features</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>↑</td>
<td>CD79B</td>
<td>CD79b</td>
<td>CD79b molecule, immunoglobulin-associated beta</td>
<td>Membrane protein that forms a heterodimer with CD79a. Together with a surface membrane immunoglobulin forms the B cell antigen receptor (BCR) complex, being the major signaling component of the receptor.</td>
</tr>
<tr>
<td>2</td>
<td>↑</td>
<td>TCL1A</td>
<td>T cell leukemia/lymphoma 1A</td>
<td>Protein involved in the phosphorylation and activation of AKT proteins that promotes nuclear translocation of AKT1. Enhances cell proliferation, stabilizes mitochondrial membrane potential, and promotes cell survival. It is located downstream of the BCR signaling pathway and more abundantly expressed in naive B cells than memory B cells. Overexpression of TCL1A prolongs naive B cell survival.</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>↑</td>
<td>HS3ST1</td>
<td>Heparan sulfate</td>
<td>Membrane protein belonging to the heparan sulfate biosynthetic enzyme family. It possesses both heparan sulfate glucosaminyl 3-O-sulfotransferase activity and anticoagulant heparan sulfate conversion activity and is a rate-limiting enzyme for synthesis of anticoagulant heparin.</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>↑</td>
<td>SH2D1B</td>
<td>SH2 domain-containing 1B</td>
<td>Single SH2-domain adapter that binds to specific tyrosine residues in the cytoplasmic tail of signaling lymphocytic activation molecule (SLAM) and related receptors. It signals downstream of CD84, which is upregulated on a major population of human memory B cells, inducing homotypic adhesion of B lymphocytes. Stimulated B cells undergo early apoptotic events in the presence of SH2D1B.</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>↑</td>
<td>MS4A1</td>
<td>Membrane-spanning</td>
<td>Membrane protein belonging to FCRL family, considered as a B lymphocyte-specific, cell-surface molecule involved in B cell activation and differentiation. Rituximab, a monoclonal antibody against the pan-B cell antigen CD20, has been successfully used in both adults and children for the management of malignant and nonmalignant immune-mediated disorders.</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>↓</td>
<td>TLR5</td>
<td>Toll-like receptor 5</td>
<td>Member of the TLR family, plays a fundamental role in pathogen recognition and activation of innate immunity by recognition of pathogen-associated molecular patterns that are expressed on infectious agents. TLR5 mediates detection of bacterial flagellins. It acts via MYD88 and TRAF6, leading to NF-κB activation, cytokine secretion, and the inflammatory response. It is highly expressed in peripheral blood leukocytes, particularly monocytes.</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>↑</td>
<td>FCRL1</td>
<td>Near 3' of Fc receptor-like 1</td>
<td>Membrane protein belonging to FCRL family, considered as a B cell coreceptor, thus involved in B cell activation and B cell differentiation. Specifically expressed by mature B lineage cells, with higher expression in naive versus memory B cells.</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>↑</td>
<td>PNOC</td>
<td>Prepronociceptin</td>
<td>Secreted protein that binds the opioid receptor-like receptor (OPRL1). It may act as a transmitter in the brain by modulating nociceptive and locomotor behavior. Altered plasma levels have been reported in patients with various pain states, depression, and liver disease. An antiinflammatory role has been reported in rat models.</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>↓</td>
<td>SLC8A1</td>
<td>Solute carrier family 8 (sodium/calcium exchanger), member 1</td>
<td>Transmembrane protein that plays a fundamental role in Ca&lt;sup&gt;2+&lt;/sup&gt;-refilling in the endoplasmic reticulum. Expressed in human macrophages and monocytes, it restores Ca&lt;sup&gt;2+&lt;/sup&gt; signals that induce TNF-α production.</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>↑</td>
<td>FCRL2</td>
<td>Fc receptor-like 2</td>
<td>Membrane protein belonging to FCRL family, expressed in spleen, peripheral blood, and bone marrow, preferentially by memory B cells.</td>
<td></td>
</tr>
</tbody>
</table>

Genes shown in boldface are B cell-related genes. Rel. expression by Tol-DF, relative gene expression — upregulation (↑) or downregulation (↓) — by Tol-DF group (median values for fold difference in gene expression for each group of the training and test sets are available in Supplemental Figure 5 and Supplemental Table 5, A and B). Annotation data are from Information Hyperlinked over Proteins (iHOP): http://www.ihop-net.org/UniPub/iHOP; NextBio: http://www.nextbio.com; and UniProt: http://www.uniprot.org/.
and impaired renal function (50% increase in their baseline CRT at the time of enrollment relative to their initial posttransplantation baseline) due to presumed immune-mediated allograft rejection. An additional group of 31 HC volunteers with no known history of renal disease/dysfunction or evidence of acute medical illness was enrolled. Group characteristics are summarized in Table 3.

Whole blood mRNA and frozen PBMCs were received by laboratories performing the selected validation assays described.

**Figure 8**
ROC curve generation combining cross-platform biomarkers. ROC curves of the training set (A) and test set (B) generated using cross-platform biomarkers and genes identified by microarray analysis. Two-class ROC curves (Tol-DF vs. nontolerant groups, excluding HCs) were generated using 4 biomarkers: B/T lymphocyte ratio, percent CD4^+^CD25^int^, ratio of anti-donor/anti-3^rd^P ELISpot frequencies, and ratio of FOXP3/MAN1A2 expression, combined with sequential addition of the 10 most significant genes. Estimated probabilities of patients from each study group of the training set (C) and test set (D) being classified as tolerant based on the cross-platform biomarker signature of tolerance (4 biomarkers plus 10 genes), calculated using a binary regression procedure.
Peripheral blood was collected in clotting positive sera were -

PBMCs were thawed on the day of the training set cohort, peripheral venous blood was drawn directly into PAXgene Blood RNA tubes (QIAGEN). Whole blood RNA was extracted using the PAXgene Blood RNA Kit including DNase I treatment (QIAGEN).

For the test set cohort, peripheral venous blood was drawn directly into Tempus Blood RNA tubes (Applied Biosystems Inc.). Whole blood RNA was extracted according to the manufacturer’s instructions (9). Total RNA samples were subjected to gene expression analysis by RT-PCR and microarrays.

Samples for RT-PCR studies. Ninety-five samples from the training set were used, consisting of 13 samples from 10 Tol-DF patients, 16 samples from 11 s-LP patients, 8 samples from 8 s-nCNI patients, 40 samples from 28 s-CNI patients, 10 samples from 9 CR patients, and 8 samples from 8 HCs. As the test set, 142 samples from the ITN cohort were used, consisting of 31 samples from 23 Tol-DF patients, 14 samples from 11 Mono patients, 52 samples from 34 s-CNI patients, 25 samples from 18 CAN patients, and 20 samples from 20 HCs.

RNA quality control. Quality and integrity of PAXgene (training set) and Tempus-purified (test set) RNA were determined using the Agilent RNA 6000 Nano Kit on the Agilent 2100 Bioanalyzer (Agilent Technologies). RNA was quantified by measuring absorbance at 260 nm on the ND-1000 Spectrophotometer (NanoDrop Technologies).

RNA amplification and labeling. Sample labeling was performed as detailed in ref. 35. Briefly, 0.5 μg total RNA was used for the amplification and labeling steps using the Agilent Low RNA Input Linear Amp Kit (Agilent Technologies) in the presence of cyamine 3-CTP. Yields of cRNA and the dye incorporation rate were measured with the ND-1000 Spectrophotometer (Thermo Scientific).

Hybridization of RISSET 2.0 Agilent custom microarrays. All whole blood samples were hybridized on the RISSET 2.0 microarray platform. This is a custom Agilent 8 × 15K 60-mer oligonucleotide microarray comprising 5,069 probes represented in triplicate. Probes selected corresponded to 4,607 genes with a valid Entrez Gene ID and an additional 407 probes that could not be assigned to a valid Entrez Gene ID. The microarray is focused on the detection of genes relevant in the field of transplantation and was designed based on current literature and published and unpublished data provided by RISSET consortium partners. Probe design was optimized for the detection of multiple transcript variants of a gene, on optimized hybridization properties of the probes, and avoiding cross-hybridization.

The hybridization procedure was performed after control of RNA quality and integrity and according to ref. 35 using the Agilent Gene Expression Hybridization Kit (Agilent Technologies). Briefly, 0.6 μg Cy3-labeled fragmented cRNA in hybridization buffer was hybridized overnight (17 hours, 65°C) to RISSET 2.0 microarrays. Following hybridization, the microarrays were washed once with Agilent Gene Expression Wash Buffer 1 for 1 minute at room temperature, followed by a second wash with preheated (37°C) Agilent Gene Expression Wash Buffer 2 containing 0.005% N-lauroylsarcosine for 1 minute. The last washing step was performed with acetonitrile for 30 seconds.

Scanning and data analysis. Fluorescence signals of the Agilent Microarrays were detected using Agilent’s Microarray Scanner System (Agilent Technologies Inc.). The Agilent Feature Extraction Software (FES version 9.5.1.1) was used to read out and process the microarray image files. To determine differential gene expression, FES-derived output data files were further analyzed using the Rosetta Resolver gene expression data analysis system (version 7.10.2.2., Rosetta Inpharmatics LLC).

First, an artificial common reference was computed from all samples included in the IOT dataset. Using this baseline, log2 ratios were calculated for the frequency of the dataset (i.e., 1 in 54,000 cells was recorded as 54,000); therefore, ratio values greater than 1.5 were defined as indicating a hyporesponsive to donor stimulation.

Blood sampling for gene expression analysis. For the training set cohort, peripheral venous blood was drawn directly into PAXgene Blood RNA tubes (QIAGEN). Whole blood RNA was extracted using the PAXgene Blood RNA Kit including DNase I treatment (QIAGEN).

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research article

for each gene and sample. Additionally, P values indicating the reliability of an observed difference between a sample and the common reference were calculated for each gene, applying the universal error model implemented in the Rosetta Resolver software (36).

Annotation enrichment analysis. Lists of genes found to be discriminatory between different sample groups, and common to both study sets, were analyzed for a statistically significant enrichment of biological pathway annotation terms in comparison to the complete RISET 2.0 microarray configuration. Term enrichment relative to the expected background distribution was scored using Fisher's exact test. Annotations were derived from different sources, e.g., Gene Ontology (GO, www.geneontology.org), signaling pathway membership, sequence motifs, chromosomal proximity, literature keywords, and cell-specific marker genes.

The data discussed in this publication have been deposited in NCBI's Gene Expression Omnibus (37) and are accessible through GEO Series accession number GSE14655 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE14655).

Quantitative RT-PCR analysis. Whole blood total RNA (200 ng) was reverse transcribed using the qQPCR First-Strand Synthesis Kit (Stratagene), and synthesized cDNA was subjected to RT-PCR analysis. Microarray data validation. A selected set of genes identified by microarray gene expression analysis were validated by qRT-PCR. qRT-PCR was performed for the following genes using pre-made TaqMan panels from Applied Biosystems: Hs01017452_m1 B lymphoid tyrosine kinase (BLK), Hs00236881_m1 CD79b molecule (CD79b), Hs01099196_m1 heparan sulfate (glucosamine) 3-O-sulfotransferase 1 (HSST1), Hs01592483_m1 SH2 domain containing 1B (SH2D1B), Hs00172040_m1 T cell leukemia (TCL1A).

Other assays screened in the training set. We also performed indirect pathway IFN-γ ELISpot, direct and indirect pathway trans-vivo delayed-type hypersensitivity assays, RT-PCR amplification for cytokine genes on direct and indirect pathway cultures of donor and recipient cells, and TCR-repertoire profiling by TCR landscape analysis.

Statistics. Nonparametric tests were used to estimate statistical significance, as n > 20 in many group comparisons and data did not conform to a normal distribution. Wilcoxon signed-rank test was used to compare responses within the same group of patients. Mann-Whitney U tests were used to compare medians between patient groups. To compare associations between clinical variables, usually recorded as categorical data and the presence or absence of anti-HLA antibodies, we used Fisher's exact test. Two-sided P values were used to indicate a significant difference when it was less than 0.05.

Statistical analysis of microarrays and biomarkers. Significantly altered expression detected by microarray was statistically determined using 4-class analysis and the Kruskal-Wallis test with Benjamini-Hochberg adjustment for FDR at 1%. We chose a nonparametric test for this analysis, as the data in some cases appeared to deviate from normality. A similar procedure was used to rank the biomarkers (tested on the log scale, with missing values set equal to the sample-wide mean).

To evaluate the predictive power of a number of variables to detect tolerant patients, we used ROC curves. To build these, first 4-class analysis identified differentially expressed probes of Tol-DF within the training set and were ranked using the Kruskal-Wallis test. Then the top-most significantly differentially expressed probes were added in a binary regression model and used to perform classification within sample. The binary regression procedure was used to compute probabilities \( p[1], \ldots, p[n] \) of being a Tol-DF patient for each subject. The ROC curve was produced by varying a probability threshold between 0 and 1; for each value of the threshold \( t \), a 2 × 2 classification table of actual class versus predicted class for subject \( i \) was set equal to “Tol-DF” if \( p[i] > t \). Bootstrap resampling of the subjects indicated that the within-sample classification results were robust. For the ITN test set, the same probes from the training set analysis were used.

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