INTRODUCTION

In a 2016 US Food and Drug Administration public meeting on patient-focused drug development, transplant recipients voiced their desire to have immunosuppressive therapy individualized and simplified to avoid side effects while ensuring efficacy. Unfortunately, when induction therapy is selected for renal transplant recipients, current evidence shows that an individual's clinical risk factors only account for 10%-33% of the observed variation in practice, while transplant center effect was responsible for the majority (51%-61%) of the variation.

Alloimmune risk stratification in renal transplantation has lacked the necessary prognostic biomarkers to personalize recipient care or optimize clinical trials. HLA molecular mismatch improves precision compared to traditional antigen mismatch but has not been studied in detail at the individual molecule level. This study evaluated 664 renal transplant recipients and correlated HLA-DR/DQ single molecule eplet mismatch with serologic, histologic, and clinical outcomes. Compared to traditional HLA-DR/DQ whole antigen mismatch, HLA-DR/DQ single molecule eplet mismatch improved the correlation with de novo donor-specific antibody development (area under the curve 0.54 vs 0.84) and allowed recipients to be stratified into low, intermediate, and high alloimmune risk categories. These risk categories were significantly correlated with primary alloimmune events including Banff ≥1A T cell–mediated rejection (P = .0006), HLA-DR/DQ de novo donor-specific antibody development (P < .0001), antibody-mediated rejection (P < .0001), as well as all-cause graft loss (P = .0012) and each of these correlations persisted in multivariate models. Thus, HLA-DR/DQ single molecule eplet mismatch may represent a precise, reproducible, and widely available prognostic biomarker that can be applied to tailor immunosuppression or design clinical trials based on individual patient risk.

KEYWORDS
clinical research/practice, clinical trial design, histocompatibility, kidney transplantation/nephrology, major histocompatibility complex (MHC), rejection: antibody-mediated (ABMR), rejection: T cell mediated (TCMR), risk assessment/risk stratification
variation. Compounding the problem, randomized controlled trials (RCTs) of immunosuppression minimization that have attempted to identify "low risk" recipients using clinical, serologic, and histologic criteria have been unsuccessful, suggesting that traditional risk factors hold little utility to personalize patient care. Thus, an unmet need in transplantation is the accurate definition of an individual's alloimmune risk for a given donor at the time of transplant: a fundamental requirement if the field is to move to precision medicine.

In 2017, the American Society of Transplantation and the American Society of Histocompatibility and Immunogenetics established the Sensitization in Transplantation: Assessment of Risk (STAR) Working Group. The goal of this expert panel is to conduct critical reviews of the pretransplant diagnostics literature and make recommendations for alloimmune risk assessment building on the 2009 Kidney Disease: Improving Global Outcomes (KDIGO) and the 2013 Transplant Society clinical practice guidelines. The STAR 2017 report created a framework that recommended 2 independent risk assessments: 1 related to the risk of immunologic memory and a second related to the risk of a primary (ie, de novo) alloimmune response posttransplant. In this context, the STAR Working Group identified that the HLA molecular mismatch was a key determinant of an individual's primary alloimmune risk and called for research to determine optimal approaches to define HLA molecular mismatch risk categories.

In this study, building on our prior work using HLAMatchmaker as a computational tool to assess donor-recipient HLA relatedness, we evaluated a novel approach to quantify HLA molecular mismatch allowing us to more precisely classify individuals into low, intermediate, or high alloimmune risk categories at the time of kidney transplant. These risk categories correlated with primary alloimmune events (ie, T cell-mediated rejection [TCMR], de novo donor-specific antibody [dnDSA] development, antibody-mediated rejection [ABMR]), as well as all-cause graft loss.

2 | CONCISE METHODS

2.1 | Study population

Approval was obtained from the institutional review board (H2011:211) and was in adherence with the declaration of Helsinki. Seven hundred twenty-four adult and pediatric consecutive renal transplants between January 1999 and July 2016 were considered for inclusion. Patients with primary nonfunction (n = 17), or pretransplant donor-specific antibody (DSA) (n = 43) were excluded, leaving 664 recipients (adult n = 606, pediatric n = 58) for analysis. Median follow-up was 91 months (range 18-226). Recipients who moved (n = 33) or died with a functioning graft (n = 112) were censored at last follow-up. Standard immunosuppression consisted of a calcineurin inhibitor (tacrolimus [87%] or cyclosporin [13%]), mycophenolate mofetil, and prednisone. Induction therapy with thymoglobulin (21%) or basiliximab (18%) was used in 39% of patients. Details on clinical, serologic, and histologic monitoring posttransplant have been reported previously and can be found in the Supplemental Methods.

2.2 | HLA typing and epitope mismatch identification

High-resolution Class II HLA typing (HLA-DRβ1/3/4/5 and HLA-DQα1/β1) was performed using sequence-specific oligonucleotide probes or sequence-specific primer technology (LABType® HD SSO, Micro SSP™; One Lambda, Los Angeles, CA). HLAMatchmaker

**FIGURE 1** HLA donor-recipient mismatch drives allore cognition. An amino acid polymorphism (yellow) present in the donor and not present in the recipient is the most basic unit of mismatch. An eplet is defined as a single polymorphic amino acid or a small patch of polymorphic amino acids within a 3 angstrom (0.3 nm) radius on or near the surface of an HLA molecule. An eplet represents the smallest functional unit of an epitope-paratope interface, which may drive antibody specificity through interactions with the central complementary determining regions of the antibody paratope. The complete epitope (green) represents all amino acids within a 15 angstrom (1.5 nm) radius typical for an antibody paratope.
software (HLA DRDQDP Matching version 2.0) was used to define Class II eplet mismatches between donors and recipients.

HLA eplet identification is based on 2 underlying principles: (a) the immune system recognizes and develops antibodies against nonself-antigens, or more specifically the epitopes on those antigens, while ignoring self-antigens/epitopes; and (b) epitope binding affinity is largely determined by a small number of polymorphic amino acids near the center of the epitope. An eplet is defined as a single polymorphic amino acid or a small patch of polymorphic amino acids within a 3 angstrom (0.3 nm) radius (Figure 1) on or near the surface of an HLA molecule. An eplet represents the smallest functional unit of an epitope-paratope interface, which may drive antibody specificity through interactions with the central complementary-determining regions of the antibody paratope. An epitope is defined by the complete antigen-antibody interface (=15 angstrom [1.5 nm]) made up of amino acids essential for specificity as well as those that affect affinity but not specificity.

2.3 | Traditional HLA mismatch vs molecular HLA mismatch assessment

Class II HLA-DR/DQ donor-recipient mismatch was evaluated by 3 different methods in this study (Figure 2). First is the traditional whole antigen method where each HLA-DRβ1 or HLA-DQβ1 donor antigen is assigned 1 mismatch if different from either recipient antigen, resulting in a sum score of 0, 1, or 2 at each locus. The second method, published previously, uses HLAMatchmaker DRDQDP (version 2) to determine the eplet mismatches for each of the HLA-DRβ1/3/4/5, HLA-DQA1, and HLA-DQB1 alleles, which are summed for each locus. In both of the first 2 methods the total score for that HLA locus is correlated with dnDSA development at that locus. The third method used HLAMatchmaker to determine the eplet mismatch for each HLA-DR or HLA-DQ molecule individually and correlated the single molecule eplet mismatch with dnDSA development against that molecule specifically (Table 1). Thresholds were then developed by receiver operating characteristic (ROC) curve analysis (see section 2.1) so that recipients could be categorized by whether any of their individual HLA-DR or DQ molecules had eplet mismatch loads above or below the thresholds. For HLA-DR each maternal and parenteral HLA-DRβ1 (n = 1328), HLA-DRβ3 (n = 481), HLA-DRβ4 (n = 392), HLA-DRβ5 (n = 231) donor alleles were considered. Donor null alleles at HLA-DRβ3/4/5 (n = 224) did not count toward the total. For HLA-DQ, α and β alleles inherited as a haplotype were considered as 1 HLA-DQA1β1 molecule (n = 1328).

2.4 | Statistics

Comparisons between baseline predictors and clinical outcomes were done using Student t test for parametric continuous variables
**TABLE 1** Comparison of HLA-DR/DQ mismatch quantification methods

| Description                                | Calculation of HLA-DR mismatch score | Calculation of HLA-DQ mismatch score | Benefits                                      | Drawbacks                                                                 |
|--------------------------------------------|-------------------------------------|-------------------------------------|-----------------------------------------------|---------------------------------------------------------------------------|
| Traditional whole antigen mismatch         | The sum of mismatched donor antigens for a given locus | DRβ1 allele 1 + DRβ1 allele 2       | 0.58                                          | Historical. Understood by all                                             |
|                                            |                                     | DQβ1 allele 1 + DQβ1 allele 2       | 0.54                                          | Imprecise. No information about the relative similarity or difference between donor and recipient antigens |
| Eplet mismatch sum                         | The sum of mismatched donor eplets for a given locus | DRβ1 allele 1 + DRβ3/4/5 allele 1 + DRβ1 allele 2 + DRβ3/4/5 allele 2 | 0.72                                          | Increased precision compared to antigen matching. One mismatch score for each HLA locus |
|                                            |                                     | DQβ1 allele 1 + DQβ3/4/5 allele 1 + DQβ1 allele 2 + DQβ3/4/5 allele 2 | 0.72                                          | Composite score of all molecules within a locus less precise in recipients with 1 antigen matched and 1 mismatched |
| Single molecule eplet mismatch             | The number of eplet mismatches for each specific molecule within each locus | DRβ1 allele 1 or DRβ1 allele 2 or DRβ3/4/5 allele 1 or DRβ3/4/5 allele 2 | 0.84                                          | Antibody specificity must be known. Specificities may overlap across molecules |
|                                            |                                     | DQβ1 allele 1 + DQβ1 allele 2 or DQβ3/4/5 allele 1 or DQβ3/4/5 allele 2 | 0.84                                          | Improved precision, particularly for recipients with 1 antigen matched and 1 mismatched |

\*AUC, area under the curve as a correlate with de novo donor-specific antibody (dnDSA) development.

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**3.1 Defining low risk for primary alloimmunity by HLA molecular mismatch**

The range of HLA-DR dz single molecule eplet mismatches was 0-22 and the range of HLA-DQ dz single molecule eplet mismatches was 0-21. Each HLA-DR dz single molecule eplet mismatch identified a molecule using the HLA-DR dz single molecule eplet mismatch thresholds, respectively. The range of HLA-DQ dz single molecule eplet mismatches was 0-31. Each HLA-DQ dz single molecule eplet mismatch identified a molecule using the HLA-DQ dz single molecule eplet mismatch thresholds, respectively.

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**RESULTS**

This consecutive cohort (n = 664) had a median follow-up of 9 months (range 18-227 months) and a median 10-year, all-cause graft survival of 74%. Screening serial sera, HLA DR dz dnDSA developed in 82 recipients (12%) at an average of 5.9 (range 0.5-17) years posttransplant. De novo DSA developed against Class I dz dnDSA had decreased graft survival compared with those who did not develop dnDSA (p < .0001). Each HLA-DR dz single molecule eplet mismatch was associated with the HLA-DQ dz single molecule eplet mismatch with an area under the curve (AUC) of 0.84 for HLA-DR dz development and 0.84 for HLA-DQ dz development, respectively (Figure S1). Recipients who developed Class II dz dnDSA alone, or Class I dz dnDSA had decreased graft survival compared with those who did not develop dnDSA (p < .0001).

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Comparisons across multiple groups were done using Kruskal-Wallis test for nonparametric data and analysis of variance for parametric test for parametric data and analysis of variance for parametric variables. Survival analysis was done by the Kaplan-Meier method, using the log-rank test for significance. ROC analysis was used to identify HLA-DR or DQ molecule specific thresholds, best associating with dnDSA development. Cox proportional hazards model was used with dnDSA development as the final model. Statistical software used was JMP (version 14.0; SAS Inc., Cary, NC).
induction therapy (P < .01) compared to Group C. Of note, risk factors known to influence dnDSA development such as nonadherence, maintenance immunosuppression, and tacrolimus coefficient of variation were similar among the 3 groups.

3.2 | Comparison of HLA-DR/DQ mismatch quantification methods

Traditional HLA-DR/DQ whole antigen mismatch, HLA-DR/DQ eplet mismatch sum thresholds (previously published HLA-DR and DQ thresholds each ≤11), and HLA-DR/DQ single molecule eplet mismatch thresholds were compared as correlates for HLA-DR/DQ dnDSA-free survival (Figure 3) and Banff ≥1A TCMR-free survival (Figure 4). Table 1 and Figures 1 and 2 outline the key differences between these 3 methods.

Traditional HLA-DR/DQ whole antigen mismatch greater than zero was associated with significantly lower HLA-DR/DQ dnDSA-free survival (P = .0003). However, there was no statistical difference in HLA-DR/DQ dnDSA-free survival between HLA-DR/DQ whole antigen risk groups other than zero (P = .48, Figure 3A). This was also true in a locus-specific analysis of HLA-DR or HLA-DQ dnDSA development (Figure S3). HLA-DR/DQ eplet mismatch sum was also associated with HLA-DR/DQ dnDSA-free survival (P = .0013, Figure 4B). Using the thresholds of ≥11 HLA-DR or DQ eplet mismatches, a trend of increased risk of Banff ≥1A TCMR across eplet mismatch groups was evident (Figure 4B). Using HLA-DR/DQ single molecule eplet mismatch risk groups, recipients with HLA-DR <7 and HLA-DQ <9 single molecule eplet mismatches (Groups A and B, n = 166, 25% of the cohort) developed Banff ≥1A TCMR in ≤10% of recipients in the first year posttransplant. However, recipients with HLA-DR ≥7 or HLA-DQ ≥9 single molecule eplet mismatches (Group C, n = 498) had significantly increased risk of Banff ≥1A TCMR (20% at 12 months, P = .0018, Figure 4C).

3.3 | Defining intermediate and high risk for primary alloimmunity by HLA molecular mismatch

Groups A and B were combined into a single Low Risk category based on the prior analysis (HLA-DR <7 and HLA-DQ <9). Because

| Group A | Group B | Group C |
|---------|---------|---------|
| DR=0 and DQ=0 | DR=1-6 and/or DQ=1-8 | DR≥7 or DQ≥9 |
| n = 93 | n = 73 | n = 498 |
| First transplant | 93% | 89% | 97% |
| Recipient age (y) | 40.4 ± 14.1 | 41.0 ± 15.0 | 44.9 ± 16.5 |
| Donor age (y) | 38.7 ± 12.8 | 40.9 ± 16.0 | 40.6 ± 15.0 |
| Living donor | 77% | 45% | 44% |
| Ethnicity (white vs other) | 77% | 62% | 65% |
| Cold ischemic time (h) | 4.2 ± 3.6 | 6.9 ± 5.6 | 7.3 ± 5.5 |
| Delayed graft function | 7% | 19% | 14% |
| Induction therapy | 20% | 27% | 44% |
| Basiliximab | 14% | 15% | 20% |
| Thymoglobulin | 7% | 12% | 24% |
| Tacrolimus vs cyclosporin | 90% | 88% | 87% |
| Tacrolimus CV 0-12 mo (n = 582) | 34.2 ± 9.4 | 39.1 ± 13.9 | 36.3 ± 12.1 |
| Mycophenolate | 100% | 100% | 100% |
| Nonadherence | 14% | 12% | 16% |

CV, coefficient of variation; ns, not significant.
Comparing HLA mismatch methods to define low risk for dnDSA. Traditional HLA-DR/DQ whole antigen mismatch (A), HLA-DQ/DQ eplet mismatch sum (B), and HLA-DR/DQ single molecule eplet mismatch (C) are correlated with de novo donor-specific antibody-free survival.
FIGURE 4. Comparing HLA mismatch methods to define low risk for TCMR. Traditional HLA-DR/DQ whole antigen mismatch (A), HLA-DQ/DQ eplet mismatch sum (B), and HLA-DR/DQ single molecule eplet mismatch (C) are correlated with Banff ≥1A T cell–mediated rejection–free survival. TCMR, T cell–mediated rejection.
Group C recipients represented 75% of the cohort, we sought to stratify these recipients further by repeating the ROC analysis at the HLA-DR and DQ loci after exclusion of recipients in the Low Risk category. For HLA-DR, no additional cutoff could be identified; however, a single molecule threshold of ≥15 HLA-DQ eplet mismatches was identified. When Group C recipients were split into groups of above or below 15 HLA-DQ eplets, 2 alloimmune risk categories were identified: Intermediate Risk (HLA-DR ≥7 and HLA-DQ ≤14, or HLA-DR 0-6 and HLA-DQ 9-14), and High Risk (HLA-DR 0-22 and HLA-DQ 15-31). Recipient demographics of the low, intermediate, and high HLA molecular mismatch risk categories are shown in Table S1. Differences between groups were found for primary transplant, recipient age, living donor, cold ischemic time, and induction therapy. There were no significant differences in baseline demographic between intermediate- and high-risk groups.

Low, Intermediate, and High HLA molecular mismatch risk categories were significantly associated with Banff ≥1A TCMR (P = .0006, Figure 5A). In a Cox model, Banff ≥1A TCMR-free survival was significantly less for recipients in the Intermediate (hazard ratio [HR] 2.02, 95% confidence interval [CI] 1.1-3.9, P = .0230) and High (HR 3.33, 95% CI 1.9-6.3, P < .0001) risk categories compared to the Low Risk category. Correlates associated with Banff ≥1A TCMR are shown in Table S2. Significant multivariate correlates of Banff ≥1A TCMR-free survival were recipient age (HR 0.98, 95% CI 0.97-0.99, P = .0037), cyclosporin vs tacrolimus (HR 5.39, 95% CI 3.6-8.0, P < .0001), delayed graft function (HR 2.04, 95% CI 1.3-3.1, P = .0035), and HLA molecular mismatch risk category (HR High vs Low 3.94, 95% CI 2.2-7.5, P < .0001; HR High vs Intermediate 1.6, 95% CI 1.1-2.4, P = .2233; HR Intermediate vs Low 2.28, 95% CI 1.2-4.5, P = .0075).

Low, Intermediate, and High HLA molecular mismatch risk categories were significantly associated with dnDSA development (P < .0001, Figure 5B). In a Cox model, dnDSA-free survival was significantly less for recipients in the Intermediate (HR 10.18, 95% CI 3.0-63.5, P < .0001) and High (HR 20.8, 95% CI 6.4-127.4, P < .0001) risk categories compared to the Low Risk category. Correlates associated with dnDSA-free survival are shown in Table S3. Significant multivariate correlates of dnDSA development were recipient age (HR 0.97, 95% CI 0.96-0.99, P = .0010), cyclosporin vs tacrolimus (HR 2.11, 95% CI 1.3-3.4, P = .0043), nonadherence (HR 2.83, 95% CI 1.7-4.6, P < .0001), and HLA molecular mismatch risk category (HR High vs Low 18.31, 95% CI 5.6-11.4, P < .0001; HR High vs Intermediate 1.84, 95% CI 1.1-3.1, P = .0149; HR Intermediate vs Low 9.96, 95% CI 2.9-62.1, P < .0001).

At least 1 renal biopsy was available in 57/72 (79%) of recipients post-dnDSA. ABMR developed in 40/57 (70%) of recipients post-dnDSA development. ABMR-free survival was significantly associated with HLA molecular mismatch risk categories (P < .0001, Figure 5C). Correlates associated with ABMR-free survival are shown in Table S4. Significant multivariate correlates of ABMR were recipient age (HR 0.98, 95% CI 0.96-0.99, P = .0116), nonadherence (HR 3.52, 95% CI 2.0-6.3, P < .0001), and HLA molecular mismatch risk category (HR High vs Low 5.06, 95% CI 2.1-14.9, P < .0001; HR High vs Intermediate 1.87, 95% CI 1.0-3.6, P = .0420; HR Intermediate vs Low 2.71, 95% CI 1.0-8.3, P = .0394).

3.4 | HLA molecular mismatch correlates with all-cause graft loss

Low, Intermediate, and High HLA molecular mismatch risk categories were significantly associated with all-cause graft loss (P = .0003, Figure S4). Correlates associated with all-cause graft loss are shown in Table S5. In a multivariate analysis, covariates significantly associated with all-cause graft loss were recipient age (HR 1.03, 95% CI 1.01-1.04, P < .0001), delayed graft function (HR 2.56, 95% CI 1.80-3.58, P < .0001), nonadherence (HR 2.00, 95% CI 1.35-2.87, P = .0006), and alloimmune risk category (HR High vs Low 1.66, 95% CI 1.12-2.53, P = .0120; HR High vs Intermediate 1.20, 95% CI 0.9-1.7, P = .3; HR Intermediate vs Low 1.39, 95% CI 0.9-1.2, P = .1).

4 | DISCUSSION

The key finding in this study is that in the absence of donor-specific memory (ie, no preformed DSA by solid phase single antigen bead assessment), quantifying the differences of HLA-DR/DQ mismatches at the molecular level can improve precision in primary alloimmune risk categorization. As a prognostic biomarker available pretransplant, its potential application includes the development of personalized immunosuppression protocols, as well as being a drug development tool for enrichment/stratification in clinical trial design to improve trial efficiency. These applications would address major unmet needs in transplantation from both the perspective of the patient and the pharmaceutical industry/academia.13,14

Traditional alloimmune risk factors reported by KDIGO and recently enumerated in the 2017 Consensus on Managing Modifiable Risk in Transplantation report were pretransplant DSA, panel reactive antibody (PRA) >0%, younger recipient age, African American ethnicity, and HLA-DR mismatch.7,15 Pretransplant DSA, a measure of alloimmune memory, correlates with ABMR, transplant glomerulopathy, and graft loss.16 However, as pretransplant DSA is avoided in most kidney transplants, methods for primary alloimmune risk stratification are needed. Although elevated PRA has been correlated with allograft outcomes, recent work using state-of-the-art antibody assessment in combination with more complete HLA typing (ie, HLA-C, HLA-DQ, and HLA-DP) has shown that when preformed DSA are ruled out, calculated PRA alone is not prognostic of graft outcomes.17-19 Younger recipient age is a well-known correlate of alloimmune risk in transplantation, likely as a result of a more robust immune system, even after adjustment for the higher prevalence of nonadherence.20 Unfortunately, there is a lack of studies to define what age cutoff may be important, and how recipient age might be used in the precision medicine context. Although certain ethnic minorities have been associated with worse outcomes, these data are confounded by socioeconomic status, HLA mismatch, and differences in drug metabolism.20 Moreover, population migration
HLA molecular mismatch category correlates with Banff ≥1A TCMR, dnDSA development, and ABMR. HLA-DR/DQ molecular mismatch categories (low, intermediate, and high) were correlated with Banff ≥1A T cell–mediated rejection–free survival (A), de novo donor–specific antibody–free survival (B), and antibody–mediated rejection–free survival (C). ABMR, antibody–mediated rejection; dnDSA, de novo donor–specific antibody; TCMR, T cell–mediated rejection.
and genetic admixture makes self-reported ethnicity increasingly imprecise such that ethnicity may be prognostic at a population level but is unlikely to have any prognostic utility at an individual level.\textsuperscript{21,22}

Since the 1950s, HLA mismatch has been known to correlate with transplant outcome.\textsuperscript{23} Unfortunately, while traditional HLA whole antigen mismatch, especially Class II, correlates with outcomes at the population level, the lack of precision limits its utility at the individual level. Traditional HLA whole antigen mismatches only evaluates whether the donor and recipient molecules are the same or different. The issue is that some mismatched HLA molecules are nearly identical while others may be very disparate—information ignored with traditional HLA mismatch assessment. Fortunately, this relative difference can be captured and quantified by HLA molecular mismatch comparisons\textsuperscript{24,25} such as the HLAMatchmaker eplet relative difference can be determined and used to categorize each recipient as low, intermediate, or high primary alloimmune risk based on the molecular mismatch analysis used in this study. Because the range of eplet mismatches is wide, it is logical to ask what cutoff might best correlate with a primary alloimmune response. De novo DSA development is a useful outcome for model development because it can be detected noninvasively, its onset can be determined by serial screening, and specificity can be assigned to the single molecule mismatch level. Previously, it was reported that the sum of adding the eplet mismatches within the same locus correlated with dnDSA development at that locus.\textsuperscript{9} However, since single molecule specificity of dnDSA can be determined in most cases, the analysis can be refined further to ask if the eplet mismatch for each individual molecule correlates with the development of dnDSA to that molecule.\textsuperscript{26} The single molecule approach would be expected to be more precise, especially for patients who have only 1 of the 2 molecules mismatched.

Using the traditional HLA whole antigen mismatch as a correlate with dnDSA development, the AUC was only 0.54 and 0.58 for HLA-DR and DQ, respectively. When applying the sum of the eplet mismatch within a locus, the AUC improved to 0.72 for both HLA-DR and DQ.\textsuperscript{9} This was enhanced further in the current study using the single molecule eplet analysis to an AUC of 0.84 for HLA-DR and DQ. Notably, results of the single molecule method provide multiple individual scores within each locus for each recipient. Thus, to bring this evaluation to the individual recipient level, thresholds were determined and used to categorize each recipient as low, intermediate, or high primary alloimmune risk based on the molecular mismatch scores across all HLA-DR\textsubscript{\(β_1/β_2/β_3/β_4/β_5\)} and HLA-DQa\textsubscript{\(α_1/α_2\)} molecules. These risk categories were highly correlated with dnDSA development, ABMR, Banff \(≤1A\) TCMR, and all-cause graft loss.

The need for reliable prognostic and predictive biomarkers at the time of transplant to allow individualization of immunosuppression in patients without alloimmune memory has been recognized.\textsuperscript{1} A prognostic biomarker is one that indicates an increased (or decreased) likelihood of a future clinical event.\textsuperscript{27} HLA-DR/DQ molecular mismatch has consistently been shown to correlate with a significantly increased risk of dnDSA development, ABMR, transplant glomerulopathy, and graft loss after adjustment for other risk factors.\textsuperscript{7-11,26,28,29} A predictive biomarker is used to identify individuals who are more likely to respond after exposure to a particular medical product or environmental agent.\textsuperscript{27} In this regard, Class II HLA eplet mismatch has been shown in 2 observational cohort studies and 1 RCT to identify high alloimmune risk patients who have increased rates of dnDSA development, rejection, and graft loss when immunosuppression is minimized through protocol-driven withdrawal or recipient nonadherence.\textsuperscript{3,9,28} Conversely, of equal importance, these studies also identified a subset of recipients with low alloimmune risk who tolerated immunosuppression minimization. If validated in prospective clinical trials, this would provide the evidence that HLA-DR/DQ single molecule mismatch can act as both a prognostic and a predictive biomarker capable of identifying which individuals require more or less immunosuppression to control their primary alloimmune response.\textsuperscript{21}

As a prognostic biomarker, HLA-DR/DQ molecular mismatch has several favorable characteristics. It is available at the time of transplant, modern HLA typing methods already provide the inputs necessary for its evaluation, and analysis software is free and is already being incorporated into HLA typing software from vendors. Thus, HLA-DR/DQ molecular mismatch score as a prognostic biomarker could significantly address this issue by enriching Phase 2 and 3 clinical trials with patients based upon risk categorization in studies evaluating novel drugs.

4.1 Limitations

Due to the relatively small sample size and the associated risk of type II error, risk quantification should be interpreted with caution, and should be validated in an independent cohort. Histology was available in 79% of recipients in the first year posttransplant; however, 97% of the death-censored graft loss occurred in the cohort with at least 1 biopsy. Methods of risk stratification will need to be tested prospectively and in independent cohorts with varying ethnicities to confirm their general applicability.

5 Conclusions

A prerequisite to precision medicine is a prognostic biomarker that correlates with clinical outcomes, is reproducible, and ideally is widely available and cost effective. Using thresholds identified for dnDSA development as an outcome, we show that the HLA-DR/DQ molecular mismatch score allows for low-, intermediate-, and high-risk stratification for Banff \(≤1A\) TCMR, dnDSA development, ABMR, and all-cause graft loss. Given that all transplant programs worldwide are supported by accredited histocompatibility laboratories,
this biomarker could be readily applied at little additional cost. Once validated, the HLA-DR/DQ molecular mismatch score could be used to tailor immunosuppression based on individual patient risk, as well as in the design of clinical trials.

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DISCLOSURE

The authors of this manuscript have conflicts of interest to disclose as described by the American Journal of Transplantation. D.N.R. is a consultant with Astellas Pharma, and P.W.N. is a consultant with Astellas Pharma and Vitaeris Inc. The other authors have no conflicts of interest to disclose.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.

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