Predicting Humoral Alloimmunity from Differences in Donor and Recipient HLA Surface Electrostatic Potential

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In transplantation, development of humoral alloimmunity against donor HLA is a major cause of organ transplant failure, but our ability to assess the immunological risk associated with a potential donor–recipient HLA combination is limited. We hypothesized that the capacity of donor HLA to induce a specific alloantibody response depends on their structural and physicochemical dissimilarity compared with recipient HLA. To test this hypothesis, we first developed a novel computational scoring system that enables quantitative assessment of surface electrostatic potential differences between donor and recipient HLA molecules at the tertiary structure level [three-dimensional electrostatic mismatch score (EMS-3D)]. We then examined humoral alloimmune responses in healthy females subjected to a standardized injection of donor lymphocytes from their male partner. This analysis showed a strong association between the EMS-3D of donor HLA and donor-specific alloantibody development; this relationship was strongest for HLA-DQ alloantigens. In the clinical transplantation setting, the immunogenic potential of HLA-DRB1 and -DQ mismatches expressed on donor kidneys, as assessed by their EMS-3D, was an independent predictor of development of donor-specific alloantibody after graft failure. Collectively, these findings demonstrate the translational potential of our approach to improve immunological risk assessment and to decrease the burden of humoral alloimmunity in organ transplantation. *The Journal of Immunology, 2018, 201: 3780–3792.

The HLA gene complex encodes highly polymorphic proteins that are the main immunological barrier to successful cell, tissue, and organ transplantation. Immune recognition of HLA class I and class II expressed on donor tissue stimulates the development of donor-specific Abs (DSA) that are the major cause of organ transplant failure in the medium to long term (1–5). Moreover, development of alloantibody through pregnancy, blood transfusion, and previous transplantation may severely limit the opportunity for organ transplantation (6, 7). Current strategies to offset the risk for development of DSA and of Ab-mediated rejection focus on minimizing the number of HLA mismatches between donor and recipient and on the administration of immunosuppression regimens that aim to suppress the recipient immune response. HLA matching is incorporated into many deceased donor organ allocation schemes, but because of the extensive polymorphism of the HLA system and the relative limitation in the size of the donor organ pool, most recipients receive allografts with one or more mismatched HLA alleles. HLA-incompatible allografts necessitate the use of increased immunosuppression, and this is a major cause of recipient morbidity and mortality (8, 9).

Current assessment of the immunological risk associated with a particular transplant is based on enumerating the number of HLA mismatches between donor and recipient and is predicated on the assumption that all mismatches within an HLA locus are of equal significance to graft outcomes. However, it is clear from animal studies that humoral alloimmunity is critically dependent on the nature of the MHC mismatch between donor and recipient, and this has been supported by observational studies in humans, suggesting that certain donor HLA are tolerated by the recipient immune system.
system (10, 11). Recent evidence shows that the potential of donor HLA to induce humoral alloresponses (HLA immunogenicity) might be a function of the number and location of amino acid sequence polymorphisms compared with recipient HLA molecules (12, 13). Numerous studies support an association between donor HLA immunogenicity, considered at the amino acid sequence level, and the likelihood of DSA developing after transplantation, and that this approach might offer superior assessment of donor–recipient histocompatibility compared with conventional HLA-matching strategies (14–17). Studies by our group have shown that the predictive ability of sequence-based HLA immunogenicity algorithms can be significantly enhanced by consideration of the physicochemical properties of amino acid polymorphisms expressed on donor HLA molecules (18–21).

Despite its promise, sequence-based assessment of HLA immunogenicity does not account for the conformational nature of antigenic recognition by BCR and for the effect of individual amino acid polymorphisms on B cell epitope structure and physicochemical properties (e.g., surface exposure, polarity, surface charge, hydrophobicity) (22–24). In particular, Ab–Ag interactions are largely governed by electrostatic forces dictated by the number and distribution of charged atoms on the surface of the HLA molecule (24–27). We have previously shown that, despite variation in their amino acid composition, HLA B cell epitopes are characterized by unique surface electrostatic potential properties that explain serological patterns of HLA-specific Ab binding (28–30). In this study, we hypothesized that the capacity of donor HLA to induce a specific alloantibody response can be predicted by quantitative assessment of their structural and surface electrostatic potential differences compared with recipient HLA molecules. We have developed a novel computational scoring system to quantify and compare HLA electrostatic properties that uses molecular modeling techniques, structural information from x-ray crystallography, and application of protein electrostatics theory. This approach was validated by analysis of HLA-specific Ab responses in a unique model of HLA sensitization comprising patients that underwent a single injection of donor lymphocytes [lymphocyte immunotherapy (LIT)] in a defined donor–recipient setting without the influence of immunosuppression or interference from other sensitizing events. The applicability of our findings in transplantation was then examined by analysis of DSA responses in patients listed for repeat renal transplantation.

Materials and Methods

Study design

The principal hypothesis that this study sought to examine is that the immunogenicity of donor HLA (defined as the potential of donor HLA to induce humoral alloimmunity in a specific recipient) can be predicted by assessment of their structural and physicochemical dissimilarity compared with recipient HLA molecules. To test this hypothesis, we developed a computational scoring system that enables quantitative assessment of surface electrostatic potential differences between donor and recipient HLA molecules at the tertiary structure level [three-dimensional (3D) electrostatic mismatch score (EMS-3D)]. The validity of the hypothesis was examined in a cohort of healthy females (recipient) that were sensitized to their male partner's lymphocytes (range 20.2–831,000 MFI). Peripheral blood was collected from all women within 2 mo before LIT and 5 wk (median: 33 d; SD: 4.5) following LIT, and serum was stored at –21°C for subsequent detection of Abs to HLA. DNA was also isolated from peripheral blood of all women and their partners for HLA typing.

This study comprised 191 consecutive women (and their male partner) who underwent their first LIT in 2009 and 2010 and had not had a previous pregnancy, blood transfusion, or organ transplantation; were not on immunosuppressive medication; and had no detectable Abs directed against their partners' HLA, as defined by complement-dependent cytotoxicity assays. During LIT, the cohort received a median (SD) of 37.4 (15.0) × 10⁶ of their partner's lymphocytes (range 20.2–831,000 MFI lymphocytes).

HLA typing

DNA samples were genotyped using the ImmunoChip, an Illumina iSelect HD custom genotyping array, according to Illumina protocols at the Institute of Molecular Biology of Kiel University. Genotype calling was performed using Illumina GenomeStudio Data Analysis software and the custom-generated cluster file of Trynka et al. (34) based on an initial clustering of 2000 UK samples with the GenTrain 2.0 algorithm and subsequent manual readjustment and quality control. Subsequent imputation of classical HLA alleles from SNP genotypes was performed using two independent HLA imputation pipelines, HLA*IMPI2 (35) and SNPHLA (36). HLA-A and -B typing was also performed (as part of the LIT protocol) using a reverse PCR sequence-specific oligonucleotide system as implemented in the Luminex platform (LABType SSO; One Lambda, Canoga Park, CA), and the results were used for quality control in case of ambiguous results. Missing genotype data from failed genotype calls or failed quality control (n = 191) were imputed, where possible, using an allele frequency-based prediction tool that considers HLA haplotype and patient race (13) (n = 39 HLA class I alleles and n = 92 HLA class II alleles), or excluded from further analysis (n = 60 HLA class II alleles).

HLA-specific Ab screening

Serum samples obtained before and after LIT were screened for HLA-specific Abs using solid-phase Luminex HLA Ab-detection beads (LABScreen; One Lambda). Selected HLA-specific Ab-positive samples were analyzed using Luminex single-Ag HLA class I and II Ab-detection beads (One Lambda). HLA single-Ag bead-defined Ab reactivity was determined using mean fluorescence intensity (MFI) cut-off thresholds of 2000 (MFI cut-off level used clinically in our center and elsewhere to define a possible alloantibody response to a given HLA) to denote the presence of DSA and of 8000 to reflect high DSA levels (widely accepted Luminex MFI level at which DSA often results in a positive donor complement-dependent cytotoxicity crossmatch test; DSA above this level commonly denotes higher immunological risk in the context of transplantation).

Comparative structure modeling of HLA alleles

Comparative structure models of all HLA class I and class II alleles represented in the HLA types of the patient cohort and of all common HLA alleles (frequency >1%) were generated using the program MODELLER via the web server (https://salilab.org/modeller/) (37). Ten comparative structure models were identified by querying the Research Collaboratory for Structural Bioinformatics Protein Data Bank (PDB) using the sequence of HLA-B*07:02 and HLA-DRB1*01:01 for HLA class I and HLA class II, respectively. The search was carried out using the Domain Enhanced Lookup Time Accelerated-Base Local Alignment Search Tool algorithm for humans (Taxonomy identification: 9606; E-value threshold of 0.005) and identified 125 HLA class II unique class I and 41 HLA class II unique class II comparative structure models. Of these, 12 HLA class I structures (PDB codes: 1K5N, 3MRE, 3CZF, 3BWA, 3LN4, 3SPV, 2BPV, 2A83, 3MBB, 1 X 7Q, 10GT, 1XH3) and 22 HLA class II structures (PDB codes: 4P4R, 3781 The Journal of Immunology
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EM-3D

The HLA type of the male partner (donor) was compared with the HLA type of the female partner (recipient) to identify mismatches in HLA-A, -B, -C, -DRB1, -DQ, and -DP loci. As shown in Fig. 1D, for HLA class I mismatches, the donor HLA was electrostatically compared with each of the recipient HLA class I alloantigens to derive the respective ESDs, and the minimum ESd value was taken to represent the EM-3D model [based on the interlocus comparison principle as previously described in our sequence-based immunogenicity algorithm (21)]. For HLA class II mismatches (Fig. 1D), the donor HLA was electrostatically compared with each of the recipient HLA within the same locus (intralocus comparison) to derive the ESDs, and the minimum value was taken to represent the EM-3D (19).

Transplant patient cohort

The patient population studied and the Ab screening protocol used have been described in detail previously (6). Briefly, the study cohort comprised 131 consecutive patients (87 males, 44 females, median age 38) who received a primary kidney allograft between 1995 and 2010 and returned to the Cambridge kidney transplant waiting list following failure of their graft during this time period (56 patients [43%] underwent transplant nephrectomy). Approximately half (50.4%) of the patients in the cohort continued to receive immunosuppression after the return to the waiting list (of those, 55% received single agent immunosuppression and 40% received dual agent immunosuppression). Ab screening was undertaken at the time of (and prior to) the first transplant after return to the transplant waiting list following graft failure and at three monthly intervals while remaining on the list for retransplantation. Screening was undertaken using Luminex single Ag beads (One Lambda), as described above. The median (SD) duration of follow up since transplantation was 2539 (1605) d.

Statistics

To investigate whether there was an association between donor HLA EM-3D and DSA development, all HLA mismatches in the patient cohort were considered together using logistic regression models. Random effects were considered to account for potential correlations within individual patients; however, these were not found to be important and not included in the final models. An MFI ≥2000 was classified as a positive result for presence of DSA and an MFI ≥8000 was classified as a high-level DSA response. Where the donor was homozygous for a particular HLA mismatch, only one observation (mismatch) was included in the model. EM-3D was modeled as a continuous variable and for illustration as a categorical variable by splitting into quartiles. To investigate HLA immunogenicity at an HLA locus and individual patient level, logistic regression models were used to examine the association between total EM-3D for HLA mismatches within a locus (one or two mismatches) and development of a recipient locus-specific DSA response, adjusting for the effect of baseline anti-donor antibody level at LIT. Median regression models were used to model the effect of EM-3D on post-LIT DSA MFI value. To assess nonlinearity of the explanatory variable, EM-3D, natural cubic spline terms were added to the logistic regression model, and these terms were kept in the model if there was sufficient evidence of nonlinearity. For analyses of the transplant cohort, logistic regression models were used to investigate the relationship between donor HLA EM-3D and DSA development at the patient level, accounting for clinical explanatory variables. Initially, each explanatory variable was modeled separately; further models investigated the additional value in incorporating EM-3D into models, including length of time to graft failure, length of time on the waiting list after listing for retransplantation, maintenance immunosuppression regimen, graft nephrectomy, and number of administered blood transfusions. Models were compared using the log likelihood ratio statistic, and p values ≤ 0.05 were considered significant. All analyses were conducted using SAS (version 9.4).

Study approval

All participating couples in the LIT cohort gave informed written consent prior to inclusion in the study for use of their data and blood samples for research, and this study was approved by the local Institutional Ethics Committee (AZ D437/09, D431/12, D474/13).

EM-3D software

We are currently developing a software suite to enable HLA model building, electrostatic potential calculation, and determination of EM-3D, which will be made freely available online (https://surgery.medschl.cam.ac.uk/divisions-and-groups/transplant-surgery/hla-structure-and-immunogenicity/immunogenicity-donor-hla-molecules/).
Results

Computational approach for calculation of HLA surface electrostatic potential and quantification of differences in electrostatic potential between HLA molecules

We generated a bioinformatics protocol to enable HLA structure prediction, surface electrostatic potential calculation, and quantification of electrostatic potential differences between different HLA molecules (Fig. 1). Because the structure of very few HLA molecules has been determined experimentally, the atomic resolution structure of a given HLA class I or class II molecule was calculated using homology modeling (MODELLER) based on information derived from high-quality HLA structures resolved by x-ray crystallography. The electrostatic potential in 3D space surrounding an HLA structure is calculated numerically by solving the linearized Poisson–Boltzmann equation for each point on a cubic grid (spacing of 0.33 Å, solvent ionic strength of 0.15 M [pH: 7.4]). Electrostatic potential comparisons consider cubic grid points within a defined region or layer of space (of thickness δ = 3 Å) at a distance σ (4 Å) above the van der Waals surface of the HLA molecule. Quantitative comparison of the electrostatic potential between two HLA molecules of interest are performed using the Hodgkin similarity index for grid points within the intersection of their layers (depicted in gray) after the two structures are superimposed, and values are converted into a distance (ESD). Derivation of EMS-3D. A mismatched donor HLA class I molecule is compared electrostatically to each of the recipient HLA class I molecules to derive the respective ESD, and the minimum ESD value (denoted by the dashed red frame) is taken to represent the EMS-3D (interlocus comparison). Similarly, for HLA class II alloantigens, the mismatched donor HLA is compared electrostatically to each of the recipient HLA within the same locus to derive the ESDs, and the minimum value is taken to represent the EMS-3D (intralocus comparison).
in an aqueous solvent, and the electrostatic potential in 3D space surrounding the HLA structure was calculated numerically by solving the linearized Poisson–Boltzmann equation (as implemented in the program APBS). After structure superimposition, comparisons of electrostatic potential between two HLA molecules of interest were performed in a defined region of space above the HLA molecular surface, and values were expressed as ESD, based on the Hodgkin index (as detailed in Materials and Methods).

The overall electrostatic potential disparity of a given donor HLA compared with recipient HLA molecules was quantified based on the EMS-3D. As described in Materials and Methods and shown in Fig. 1D, for HLA class I alloantigens, the mismatched donor HLA was compared electrostatically to each of the recipient HLA class I molecules to derive the respective ESDs, and the minimum ESD value was taken to represent the EMS-3D [intralocus comparison (21)].

Similarly, for HLA class II alloantigens, the mismatched donor HLA was compared electrostatically to each of the recipient HLA within the same locus to derive the ESDs, and the minimum value was taken to represent the EMS-3D [intralocus comparison (19)].

**Amino acid sequence polymorphism and disparities in surface electrostatic potential among HLA class I and class II alleles**

To investigate the relationship between amino acid sequence polymorphisms among HLA molecules and differences in their surface electrostatic potential, we performed pairwise, all-versus-all comparisons between common (frequency >1%) HLA alleles within individual HLA class I and class II loci. The ESD between HLA ranged from 0.00 to 0.777 [median: 0.307; interquartile range (IQR): 0.219–0.379], reflecting the overall structural and physicochemical similarity of molecules within the same protein family (Supplemental Table I). Overall, there was poor correlation between amino acid sequence polymorphism and electrostatic disparity for compared HLA class I (R² = 0.439) and class II molecules (R² = 0.317) with wide variation of ESD values for the same level of sequence polymorphism (Supplemental Fig. 1). It was notable that disparities in electrostatic potential were highest among HLA-DQ alloantigens and lowest among HLA-DR and -DP alloantigens, whereas HLA class I molecules had similar levels of variation in their electrostatic properties. Supplemental Fig. 2 shows the ESDs for pairs of compared HLA alleles within individual HLA loci presented as symmetrical heat maps with reordering such that electrostatically similar alleles are clustered together.

**LIT as a model of humoral alloimmunity**

Investigation of alloantibody responses against donor HLA in human transplantation is commonly confounded by many, often difficult to control, factors such as differences in allosensitization events (e.g., etiology [pregnancy, transfusion of blood products, and/or previous transplant], number, and time point), in disease context, and in immunosuppression regimens within the examined patient cohort. We overcame these limitations by studying HLA-specific alloantibody development in a unique patient cohort comprising healthy females that received a single intradermal injection of PBLs obtained from their partner as part of their treatment for infertility with LIT. The cohort comprised 191 couples with a median (SD) age of 34 (3) for females and 37 (4) for males. Comparison of male (henceforth referred to as donor) and female (henceforth referred to as recipient) HLA types revealed that the patient cohort was highly mismatched with a median number of 8 (IQR: 6–9) out of possible 12 HLA class I and class II mismatches (Fig. 2). HLA mismatches between individual donor–recipient pairs were pooled and analyzed together, accounting for potential effects at the patient level (see Materials and Methods).

After exclusion of HLA alleles that could not be determined at two-field level (donor HLA alleles n = 38; recipient HLA alleles n = 22), HLA mismatches where preformed DSA was identified on pre-LIT Ab screening with Luminex solid-phase assays (n = 9) and HLA mismatches that were not represented in the Luminex single-Ag bead panel (n = 176), 1381 HLA mismatches were considered for further analyses (242 HLA-A, 266 HLA-B, 213 HLA-C, 257 HLA-DRB1, 247 HLA-DQ, and 156 HLA-DP).

**FIGURE 2.** Distribution of HLA mismatches in the LIT patient cohort. This figure shows the percentage of patients (lymphocyte donors) with 0, 1, or 2 mismatches within HLA-A, -B, -DRB1, -DQ, and -DP loci.

**HLA-specific alloantibody responses after LIT**

Following LIT, HLA-specific Ab detection using Luminex showed that Ab binding against mismatched donor HLA had a median (IQR) MFI of 1039 (42, 5548). The majority of immunized recipients (84%) developed an IgG DSA response (defined as MFI ≥2000) against one or more mismatched HLA expressed on their partner’s lymphocytes. Overall, DSA was detected against 569 of the 1381 (41%) donor–recipient HLA mismatches with a median (IQR) MFI of 6939 (3795, 9917). Luminex-detected binding against donor HLA-C and -DP mismatches was of low magnitude (median [SD] MFI of 40.0 [1115.0] for HLA-C and 102.4 [2868.2] for HLA-DP), and DSA responses were less frequent (DSA against 12 of 213 [6%] donor HLA-C mismatches and against 32 of 156 [21%] donor HLA-DP mismatches) compared with other loci (Fig. 3). Strong DSA responses (in frequency and magnitude) were noted against mismatches at HLA-A (160 of 242, 66%; median [IQR] MFI: 7610 [5406, 10295]), -B (138 of 266, 52%; median [IQR] MFI: 6852 [3546, 9353]), and -DQ (136 of 247, 55%; median [IQR] MFI: 7610 [5406, 10295]) loci, followed by development of DSA against donor HLA-DRB1 (91 of 257, 35%; median [IQR] MFI: 4493 [3466, 9264]) alloantigens (Fig. 3).

**Donor HLA immunogenicity and risk of development of DSA**

We examined the association between development of DSA and the immunogenicity of donor HLA mismatches as determined by comparative assessment of electrostatic potential between donor and recipient HLA (EMS-3D). Fig. 4 shows the frequency of donor–recipient HLA mismatches according to their EMS-3D grouped by HLA locus; overall, the median (IQR) EMS-3D was 0.30 (0.24–0.35) and 0.22 (0.18–0.32) for HLA class I and class II, respectively. DSA responses against HLA-C mismatches were infrequent (Supplemental Fig. 3), reflecting the relatively low expression of HLA-C on lymphocytes (52), and were therefore not
Donor homozygosity for a given HLA class I or II mismatch had no effect on the risk of DSA development (data not shown). Although there was evidence that alloantibody responses were more likely the higher the amount of donor lymphocytes administered during LIT (adjusted OR: 1.02 per 10^6 increase in lymphocyte dose, 95% CI: 1.01–1.03; \( p < 0.0001 \)), adjusting for donor lymphocyte dose did not alter the relationship between donor HLA EMS-3D and risk of DSA development. In the analyses above, donor–recipient HLA mismatches and their immunogenic potential were considered individually and independently from each other. To consider HLA immunogenicity at a locus and individual patient level, we examined the association between the overall immunogenic potential of HLA mismatches within a locus (as assessed by EMS-3D) and the likelihood of a recipient locus-specific DSA response [as suggested by other authors (14, 63)]. This analysis showed that locus-specific EMS-3D was strongly associated with HLA-A, -B, -DRB1, and -DP DSA development (OR: 1.40 per 0.1 U increase; 95% CI: 1.21–1.61; \( p < 0.0001 \)) for donor HLA-A and -B; OR: 1.57 per 0.1 U increase; 95% CI: 1.40–1.77, \( p < 0.0001 \) for HLA-DR, -DQ, and -DP administered during LIT. As shown in Fig. 5, this analysis demonstrated a strong relationship between donor–recipient HLA electrostatic disparity and predicted probability of an alloantibody response for all loci examined. Wider CIs were observed for the immunogenic potential of high EMS-3D HLA-DR and -DP alloantigens, and this reflected the relatively low number of observations for such mismatches in the patient cohort. The association was strongest for HLA-DQ alloantigens (Fig. 5D), and it is notable that multiple recent studies have highlighted the predominance of HLA-DQ–specific humoral alloresponses after solid organ transplantation (57–60). Overall, our model predicts that donor HLA class I and class II (HLA-A, -B, -DRB1, -DQ, and -DP) with low EMS-3D have an ~10% probability of inducing DSA (e.g., the observed probability of an alloantibody response for HLA with EMS-3D <0.045 was 11% [of 27 mismatched HLA-3–induced DSA]). This probability increases to over 75% for donor HLA with the highest EMS-3D (e.g., the observed probability of a DSA response for donor HLA with EMS-3D >0.38 was 71% [of 110 mismatched HLA–induced DSA]). Fitting the model to examine high-level DSA responses (defined as DSA MFI ≥8000, which, in the context of transplantation, commonly denotes high immunological risk) showed a near linear association between donor HLA EMS-3D and predicted probability of DSA development (Fig. 5G).

Relationship between donor HLA EMS-3D and DSA MFI level after LIT

We next considered the relationship between donor HLA EMS-3D and the magnitude of the alloantibody response as assessed based on the MFI binding detected in the Luminex assay. The latter provides semiquantitative information on the level of circulating alloantibody, and previous studies have shown an association between DSA MFI level and clinical outcome (5, 61, 62). Median regression analysis showed that donor HLA with increasing EMS-3D were associated with progressively stronger (higher MFI) alloantibody responses following LIT (\( p < 0.001 \); Fig. 6). The magnitude of the alloantibody response increased from a median MFI of 48 (IQR: 0–300) for donor HLA class I and II (HLA-A, -B, -DRB1, and -DQ) with EMS-3D <0.14 to a median MFI of 6432 (IQR: 1,876–10,002) for alloantigens with EMS-3D >0.35 (Fig. 6). The association between donor HLA EMS-3D and MFI binding level was strongest for donor HLA class II mismatches (Fig. 6B).

Donor homozygosity for a given HLA class I or II mismatch had no effect on the risk of DSA development (data not shown). Although there was evidence that alloantibody responses were more likely the higher the amount of donor lymphocytes administered during LIT (adjusted OR: 1.02 per 10^6 increase in lymphocyte dose, 95% CI: 1.01–1.03; \( p < 0.0001 \)), adjusting for donor lymphocyte dose did not alter the relationship between donor HLA EMS-3D and risk of DSA development. In the analyses above, donor–recipient HLA mismatches and their immunogenic potential were considered individually and independently from each other. To consider HLA immunogenicity at a locus and individual patient level, we examined the association between the overall immunogenic potential of HLA mismatches within a locus (as assessed by EMS-3D) and the likelihood of a recipient locus-specific DSA response [as suggested by other authors (14, 63)]. This analysis showed that locus-specific EMS-3D was strongly associated with HLA-A, -B, -DRB1, and -DP DSA development (OR: 1.40 per 0.1 U increase; 95% CI: 1.21–1.61, \( p < 0.0001 \) for HLA-A and -B; OR: 1.57 per 0.1 U increase; 95% CI: 1.40–1.77, \( p < 0.0001 \) for HLA-DR, -DQ, and -DP) independent of lymphocyte dose administered during LIT.
FIGURE 4. Frequency of HLA class I and class II mismatches in the LIT patient cohort according to their EMS-3D. The figure depicts the frequency of donor-recipient HLA mismatches according to their EMS-3D, grouped by HLA locus. The median (IQR) EMS-3D for individual loci was HLA-A: 0.32 (0.27–0.36); HLA-B: 0.28 (0.22–0.33); HLA-C: 0.32 (0.22–0.40); HLA-DRB1: 0.20 (0.17–0.24); HLA-DQ: 0.35 (0.20–0.42); and HLA-DP: 0.19 (0.18–0.24).
natural cubic splines were fitted.

HLA-DRB1; OR: 1.90 per 0.1 U increase in EMS-3D, 95% CI: 1.07–3.23, p = 0.0026 for HLA-DQ).

OR on Developing HLA DSA (MFI ≥ 2000)

| HLA Locus | Donor HLA MM, DSA | OR (95% CI) | p Value |
|-----------|------------------|-------------|---------|
| HLA-A     | n = 242, 160 events | 1.64 (1.14, 2.37) | 0.007 |
| EMS-3D    | n = 266, 138 events | 1.57 (1.15, 2.15) | 0.004 |
| HLA-B     | n = 257, 91 events | 2.64 (2.01, 3.48) | <0.0001 |
| EMS-3D    | n = 156, 32 events | 3.32 (1.41, 7.81) | 0.004 |
| HLA-DRB1  | n = 508, 298 events | 1.70 (1.35, 2.15) | <0.0001 |
| EMS3D     | n = 660, 259 events | 2.56 (2.13, 3.07) | <0.0001 |
| HLA-DQ    | n = 1168, 557 events | 2.35 (2.04, 2.71) | <0.0001 |

Logistic regression models were used to investigate the association between donor HLA EMS-3D and donor-specific alloantibody development. ORs are omitted where natural cubic splines were fitted.

**Analysis of HLA-specific Ab responses after kidney transplantation**

We next considered, in a proof of principle study, the applicability of our approach in the kidney transplantation setting. Alloantibody responses against donor HLA expressed on renal allografts were examined in a cohort of 131 kidney transplant recipients returning to the transplant waiting list following first graft failure. Humoral responses against HLA class I alloantigens predominate long term after kidney transplantation and are strongly associated with graft failure (1), and, therefore, this analysis focused on development of DSA against mismatched HLA class II alloantigens. The demographic and transplant characteristics of the patient cohort have been published previously (6). To account for factors that may influence HLA-specific Ab responses at an individual patient level, multivariable logistic regression analysis of the association between EMS-3D of donor kidney HLA-DRB1 and -DQ mismatches and DSA development was adjusted for length of time to graft failure, length of time on the waiting list after listing for retransplantation, maintenance immunosuppression regimen while on the transplant waiting list, graft nephrectomy, and number of administered blood transfusions. Similar to the findings on DSA responses after LIT, this analysis showed that the EMS-3D of HLA-DRB1 and -DQ mismatches expressed on donor kidneys were independently correlated with the risk of development of DSA after kidney transplant failure (OR: 1.86 per 0.1 U increase in EMS-3D, 95% CI: 1.07–3.23, p = 0.028 for HLA-DRB1; OR: 1.90 per 0.1 U increase in EMS-3D, 95% CI: 1.25–2.88, p = 0.0026 for HLA-DQ).

**Discussion**

The capacity of donor HLA to stimulate alloantibody responses (HLA immunogenicity) is dependent upon their structural recognition by receptors on recipient B cells that initiate the immune response, and previous work has suggested that HLA immunogenicity should be considered in the context of amino acid sequence polymorphisms between donor and recipient HLA molecules (12, 21, 64). The present investigation introduces a fundamentally new approach at evaluating the immunogenic potential of donor HLA focusing entirely on their tertiary structure and on their unique structural and surface electrostatic potential properties compared with recipient HLA molecules. We have developed a computational approach to compare and quantify HLA electrostatic properties at atomic resolution level and applied it to predict HLA-specific alloantibody development in a unique model of human sensitization. We show that 1) HLA molecules differ widely at the level of electrostatic potential in 3D space, and these differences are not explicable on account of the underlying amino acid sequence polymorphisms; 2) the electrostatic disparity of a donor HLA compared with recipient HLA molecules, as assessed by EMS-3D, was strongly associated with the risk of development of donor-specific alloantibody; and 3) electrostatic potential disparities are highest among HLA-DQ molecules, which were the most immunogenic alloantigens in this study and whose immunogenicity conform best to our EMS-3D algorithm. Taken together with our proof of principle study in the setting of human kidney transplantation, the present investigation provides, to our knowledge, important first validation that donor HLA immunogenicity can be predicted based on assessment of their unique surface electrostatic potential properties compared with recipient HLA molecules.

The risk of allosensitization after transplantation increases incrementally with the number of HLA mismatches at individual HLA class I and class II loci (6). However, simple enumeration of differences at the whole Ag level is constrained by limited possible values (zero, one, or two mismatches per locus) and does not account for differences in donor HLA immunogenicity for a given recipient. Current approaches for determining the potential of a donor HLA to induce an alloantibody response are based on quantifying the degree of dissimilarity between the donor and recipient HLA molecules (12, 17, 19, 21, 65). The most frequently used methods (HLAMatchmaker and Cambridge HLA immunogenicity algorithm) evaluate differences in the number and location of amino acid mismatches at continuous and discontinuous (eplets) positions on the HLA sequence, and multiple studies have suggested they provide superior risk stratification over conventional HLA mismatch grade for predicting development of DSA, allograft rejection, transplant glomerulopathy, and allograft survival (14, 16, 18, 20, 63). Both of these methods reflect differences in amino acid sequence between donor and recipient HLA mismatches and generate highly correlated scores that provide a similar assessment of HLA immunogenicity (18, 66). Importantly, accounting for the physicochemical properties of donor HLA amino acid polymorphisms appears to improve prediction of DSA development against HLA class I alloantigens (18, 19).
FIGURE 5. Probability of donor-specific alloantibody response after LIT according to the EMS-3D of mismatched HLA on donor lymphocytes. The relationship between the immunogenic potential of donor HLA, as determined by EMS-3D, and the probability of a donor-specific alloantibody response after LIT was examined using logistic regression modeling. Each panel shows a logistic regression model with 95% CI (dotted lines) for individual HLA loci (A–E) and for HLA class I and class II loci combined (F and G). DSA responses against HLA-C mismatches were infrequent and were not examined. Donor-specific alloantibody responses were defined using MFI cut-off thresholds $\geq 2000$ (A–F) and $\geq 8000$ (G). Wide CIs for alloantibody responses against HLA-DR and -DP alloantigens reflect the relatively low number of observations for HLA-DR and -DP mismatches with high EMS-3D scores in the LIT patient cohort. Relatively few alloantibody responses with MFI $\geq 8000$ were noted against HLA-DP alloantigens ($n = 7$), and, therefore, HLA-DP mismatches were not included in the (G) model.
Protein electrostatics reflect the amino acid composition of their primary structure but are mainly determined by the number and distribution of polar and charged residues, the protonation state of ionizable groups within a given ionic environment, and their ability to form specific bonding interactions, such as salt bridges and hydrogen bonds. Importantly, our study shows that the variation in surface electrostatic potential between HLA molecules cannot be inferred on account of differences at the amino acid sequence level, and relatively poor correlation exists between residue polymorphisms and electrostatic disparity among HLA class I and class II molecules (Supplemental Fig. 1). Given that hydrophobic patches on a protein surface tend to have low electrostatic potential compared with an acidic, basic or polar patch, the electrostatic potential also captures aspects of the protein’s hydrophobic interaction properties. Electrostatic forces are important determinants of the affinity and specificity of macromolecular interactions, and it has been suggested that the process of affinity maturation involves optimization of electrostatic interactions in the BCR–Ag binding site (24, 25, 67, 68). Our study showed that donor HLA with high versus low EMS-3D were more likely to induce a specific alloantibody response, and, therefore, it would be tempting to speculate that alloantigens with disparate electrostatic potential profiles compared with recipient HLA molecules lead to more efficient BCR recognition in the secondary lymphoid organs and to improved selection and survival of differentiated B cells during the process of affinity maturation in the germinal center. Indeed, recent insights into the mechanisms that determine the fate decision of proliferating, Ag-activated B cells at the pregerminal center stage suggested that B cells with higher affinity to their Ag presented more HLA peptide to and made longer-lasting contact with cognate T follicular helper cells at the B cell–T cell border in secondary lymphoid organs, resulting in more T cell help and differentiation into germinal center B cells (with further BCR diversification through somatic hypermutation) (69, 70). In contrast, proliferating B cells with lower affinity to their Ag may form less durable T follicular helper cell–B cell conjugates and are more likely to develop into germinal center–independent memory B cells (that undergo class switching but not somatic hypermutation) (71). The implication of this model of Ag-activated B cell differentiation for the present investigation is that alloantibody responses to donor HLA with high EMS-3D might be derived by germinal center–dependent B cells and are of high affinity, whereas humoral responses to donor HLA with lower EMS-3D, when triggered, might be derived by germinal center–independent B cells and are more broadly reactive and of lower affinity. It would be interesting to investigate this hypothesis in future studies.

Multiple studies over recent years have provided strong evidence in support of the association between the development of donor HLA-specific alloantibodies and the risk of acute Ab-mediated rejection, chronic rejection, and allograft loss across all solid organ transplants (63, 72–77). The ability to assess the risk of posttransplant humoral alloimmunity associated with particular donor–recipient HLA combinations is of major clinical interest, both to inform organ allocation policies and to enable more efficient immune monitoring and individualization of immunosuppression protocols to help prevent de novo DSA development.
The present study suggests that the probability of an alloantibody response (generation and magnitude) against a donor HLA-A, -B, -DRB1, -DQ, or -DP alloantigen increases with increasing EMS-3D. Although our study does not enable identification of an HLA immunogenicity threshold in the setting of clinical transplantation, our findings suggest that it might be possible to identify a substantial number of low EMS-3D HLA mismatches that might be tolerated by the immune system of a given recipient. DSA development against both HLA class I and class II alloantigens increases the risk of subsequent rejection and allograft failure, but humoral responses against HLA class II seem to predominate, and these most commonly involve HLA-DQ–specific alloantibodies (58–60, 78). Our analysis of the most common HLA-DQ alleles (Supplemental Fig. 1) showed that electrostatic potential disparities are highest among HLA-DQ alloantigens (with only a modest correlation between ESD and the underlying amino acid sequence polymorphism) compared with other loci, and this may account for their increased immunogenic potential. Notably, HLA-DQ immunogenicity conformed best to our prediction model with a strong association between donor EMS-3D and probability of a DSA response in the LIT cohort, whereas a strong association between HLA-DQ EMS-3D and DSA development was also noted in the kidney transplant cohort.

The present study has focused on the structural aspects of donor HLA allorecognition that influence the subsequent humoral response by recipient B cells. Our computational protocol enables quantification of electrostatic potential differences between donor and recipient HLA, accounting for the entire 3D space around the HLA molecule to produce an average score. Exclusion of the membrane-bound area of the HLA did not alter the results of our analysis as this part of the molecule is relatively monomorph and therefore similar among different HLA. However, it is possible that a small surface area on a donor HLA that differs widely in electrostatic potential from the respective area on recipient HLA might be sufficient to drive the alloimmune response, although the average difference across the entire molecule remains low. Our computational method can be adapted to incorporate immunogenic “hot-spots” on the HLA molecular surface (e.g., functional B cell epitopes), and this is the subject of our current research (28). It is also important to recognize that proliferation and differentiation of Ag-specific naive B cells into memory B cells and long-lived plasma cells requires T cell help through linked recognition of Ag-derived peptides presented in the context of B cell HLA class II molecules. Previous observational studies highlighted the importance of the HLA-DR phenotype of the recipient in humoral alloresponses to donor HLA class I alloantigens (79, 80), and, more recently, this concept has been extended to evaluate the capacity of recipient HLA-DR molecules to bind donor HLA class I– and class II–derived peptides using the NetMHCIIpan computational method and to examine the contribution of this pathway to DSA development (81–84). Computational prediction of HLA class II–restricted epitopes by CD4+ T cells is of great interest for understanding immune responses in the context of transplantation, autoimmunity, infection, and cancer, but it is a difficult and complex undertaking because of the open conformation of the HLA class II peptide binding groove that can accommodate peptides of variable length (10–30 aa long) and because of the fact that Ag processing and peptide loading are incompletely understood (85). Nevertheless, this is the subject of intense research, and a multitude of computational methods are currently available for predicting HLA class II peptide binding, with mixed results (86, 87). Despite the strong association between the surface electrostatic potential properties of donor HLA and alloantibody responses in the current study, DSA development was noted against HLA with low EMS-3D and vice versa. Consideration of HLA class II peptide presentation to CD4+ T cells is likely to improve the predictive ability of our immunogenicity algorithm, and we are currently undertaking relevant studies to explore this question. To this extent, it would be intriguing to examine the contribution of electrostatic interactions within the pockets of the HLA class II peptide binding groove to high-affinity peptide binding (49, 88).

A strength of the current study is that HLA immunogenicity was investigated in a unique model of HLA sensitization comprising nonsensitized individuals that underwent a single sensitizing event (injection of donor lymphocytes), followed by detection of HLA-specific development approximately 5 wk later. To our knowledge, this is the first study to systematically examine HLA immunogenicity in this setting, thus avoiding commonly encountered confounders in similar studies, including variations in baseline disease, nonuniform immunosuppression protocols, and unknown and variable sensitization events (previous transplants, pregnancies, and blood transfusions). We acknowledge that it would have been interesting to examine the temporal evolution of the immune response at different time points, but this was not possible as further serum samples were not routinely collected. The kinetics of IgG HLA-specific Ab development after exposure to human lymphocytes have not been studied in detail, especially using sensitive detection methods, but it is well documented that specific humoral responses (class switched and affinity matured) peak within 30 d from exposure to Ag (71), and a previous study in the context of LIT suggested that the alloresponse peaks during the second month after lymphocyte immunization (89). It is, therefore, unlikely that relevant alloantibody responses have been missed because of the timing of serum collection in this study. Another limitation of the study is that a relatively small cohort of patients was examined to investigate development of DSA after failure of kidney transplant, and our findings would be strengthened if they were confirmed in larger cohorts and in patients with functioning grafts that are prospectively monitored for alloantibody development, accounting for immunosuppression regimen, drug levels, and for noncompliance. Similarly, our study was not powered to compare the predictive ability of our structural HLA immunogenicity algorithm relative to currently available, amino acid sequence–based methods, and, therefore, future studies are warranted to address this limitation and help identify the best approach, or combination of methods, to improve immunological risk assessment in the clinical transplantation setting. We were unable to systematically assess the immunogenicity of HLA-C and -DP alloantigens because of their relatively low expression on lymphocytes. Alloantibodies against HLA-DP can cause humoral rejection after kidney transplantation, and mismatching at the -DP locus is associated with decreased graft survival in patients undergoing repeat kidney transplantation (90, 91). Within the confines of the lymphocyte HLA sensitization model, HLA-DP immunogenicity seemed to conform to our algorithm, and it would be interesting to examine the applicability of our approach in the transplant setting, both in terms of predicting HLA-DP–specific sensitization and for analysis of the relevant effect of -DP mismatching on renal transplant outcomes. Finally, this study focused on DSA detection based on the Luminex solid phase assay. It would be interesting to examine a more functional readout of the humoral response, such as the ability of DSA to cause complement-dependent cell lysis. However, the latter would require access to very large panels of lymphocytes with appropriately selected HLA types and would need careful interpretation of the results (e.g., levels of HLA expression among different lymphocyte panels, potential cross-reactivity with multiple HLA targets expressed on lymphocytes, etc.). Alternatively, the Luminex C1q assay has been used to detect complement-binding alloantibodies. We, and others, have
questioned the clinical value of the LumineX C1q assay readout and showed that C1q binding in the LumineX assay is closely related to Ab MFI level (92–94). Accordingly, based on our finding that donor HLA EMS-3D correlated strongly with DSA defined at MFI >8000, we would anticipate a similarly strong relationship between EMS-3D and C1q binding DSA. Such an analysis, however, was beyond the scope of our study.

In conclusion, the present investigation demonstrates a clear relationship between the electrostatic properties of HLA molecules and their immunogenetic potential. Quantification of electrostatic potential differences at the tertiary level between donor and recipient HLA molecules enables prediction of humoral allosestic responses in the context of lymphocyte allosensitization. We have shown the translational potential of this approach in the clinical setting of kidney transplantation. Our approach has the potential to identify acceptable HLA mismatches for a given recipient, thereby increasing access to suitable donors that are currently considered a poor match, to enable better immunological risk assessment, and to decrease the burden of allosensitization and humoral rejection after solid organ transplantation.

Disclosures

The authors have no financial conflicts of interest.

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