Impact of HLA Diversity on Donor Selection in Organ and Stem Cell Transplantation

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Abstract

The human major histocompatibility complex is a multigene system encoding polymorphic human leucocyte antigens (HLA) that present peptides derived from pathogens to the immune system. The high diversity of HLA alleles and haplotypes in the worldwide populations represents a major barrier to organ and allogeneic hematopoietic stem cell transplantation, because HLA incompatibilities are efficiently recognized by T and B lymphocytes. In organ transplantation, pre-transplant anti-HLA antibodies need to be taken into account for organ allocation. Although HLA-incompatible transplants can be performed thanks to immunosuppressive drugs, the de novo production of anti-HLA antibodies still represents a major cause of graft failure. The HLAMatchmaker computer algorithm determines the immunogenicity of HLA mismatches and allows to define HLA antigens that will not induce an antibody response. Because of the much higher stringency of HLA compatibility criteria in stem cell transplantation, the best donor is a HLA genotypically identical sibling. However, more than 50% of the transplants are now performed with hematopoietic stem cells from volunteer donors selected from the international registry. The development of European national registries covering populations with different HLA haplotype frequencies is essential for optimizing donor search algorithms and providing the best chance for European patients to find a fully compatible donor.

Key Words
HLA polymorphism · Kidney transplantation · Hematopoietic stem cell transplantation · Haplotype frequency · Alloreactivity · Epitopes

Introduction

Human leucocyte antigens (HLA) are encoded in the major histocompatibility complex (MHC) and represent the most polymorphic genes in all human populations. Although originally defined as blood groups present on leucocytes, HLA molecules are expressed on the surface of virtually all nucleated cells. Their biological role is to present peptides derived from pathogens or altered self-proteins to the immune system of the host. A cellular and humoral adaptive immune response is initiated after the recognition of HLA-peptide complexes by specific recep-
tors on T lymphocytes. This defence mechanism is highly efficient in fighting against infections but constitutes a major barrier to transplantation because T and B cells are capable to recognize allogeneic (i.e. incompatible) HLA molecules and develop a cellular and/or humoral response to mismatched HLA antigens. Host-pathogen interactions have contributed to the selection of HLA-compatible donors in organ and hematopoietic stem cell transplantation (HSCT). The genetic variability within a population and also between different populations has therefore to be considered in organ exchange programs and in unrelated hematopoietic stem cell (HSC) donor search algorithms.

**HLA Polymorphism**

The human MHC on the short arm of chromosome 6 comprises HLA class I (A, B, C) and class II genes (DRA, DRB1, DRB3, DRB4, DRB5, DQA1, DQB1, DPA1, DPB1; fig. 1) [1]. HLA loci encode cell surface HLA molecules (fig. 1) which are characterized by a high degree of polymorphism. In clinical laboratories, about 100 different HLA serotypes (A, B, C, DR, DQ) can be defined on peripheral blood lymphocytes using a complement-dependent cytotoxicity assay with HLA-specific alloantisera. Since the cloning and sequencing of the first HLA class I (A, B) and II genes (DRA/DRB1) in the early 1980s, the number of new HLA variants has increased continuously. Virtually all serotypes are now subdivided into several alleles (2 to >400) leading to more than 9,000 different alleles in the worldwide populations, each assigned a given name by the WHO HLA Nomenclature Committee [2] (www.ebi.ac.uk/ipd/imgt/hla). Although most of the polymorphisms are located in the exons encoding the very distal part of HLA molecules (α1/α2 domains for class I antigens and α1/β1 domains for class II antigens; fig. 1) forming the peptide-binding cleft, variations are also located elsewhere in the genes. About 25% of all HLA class I and II alleles differ by silent substitutions, and 3% are characterized by mutations in coding or non-coding regions that lead to a lack of cell surface expression (null alleles; fig. 1).

The HLA alleles and haplotypes present large variations in frequencies within the worldwide populations [3–6]. Even within a given country, regional differences can be detected [7, 8]. Because of the enormous diversity of allelic combinations, no more than 20 HLA-A,B,DRB1 haplotypes occur at a frequency of more than 0.01 [4]. A recent international survey of sequenced alleles allowed to estimate that 14.3% of all HLA class I and II alleles (IMGT/HLA Database release 3.9.0) occurred at a frequency of more than 0.0001 or were identified in ≥5 unrelated individuals or in ≥3 individuals with a specific haplotype [9]. This means that the large majority of alleles are found at extremely low frequencies and should not significantly impact patient-donor matching in clinical transplantation.

Combinations of HLA antigens encoded by genes on the same chromosome are referred to as HLA haplotypes. A Mendelian segregation of HLA-A,B,DRB1 haplotypes is illustrated in figure 2. The probability that a sibling shares the same two haplotypes with a patient is 25%. The probability of sharing only one haplotype is 50%. In that case, the sibling is said to be haplodenital and, by
definition, each parent is also haplIDENTical to the pa-

tient. In the example shown in figure 2, segregation of the

haplotypes can be defined unambiguously on the basis of HLA-A,B,DRB1 typing. Because HLA-C is located between the A and B loci and because the DRB1 and DQB1 loci are closely linked, A,B,DRB1 matching implies C and DQB1 matching. HLA-DPB1 matching is expected in about 98% of the cases, taking into account a 1–2% recombin-

ation rate between DQB1 and DPB1 loci. F = Father; M = mother;
P = patient; S1 = HLA-identical sibling 1; S2 = HLA-haplIDENTical

sibling 2; S3 = HLA-incompatible sibling 3. For simplification, the
genotypes are indicated using the serologic nomenclature.

**Fig. 2.** Mendelian inheritance of HLA-A,B,DRB1 haplotypes in a
family study. A patient and a sibling donor have a 25% chance to
be genotypically HLA identical, i.e. sharing the same 2 haplotypes,
and a 50% chance of being HLA haplIDENTical, i.e. sharing one
haplotype. In most cases the segregation of the haplotypes can be
defined unambiguously on the basis of HLA-A,B,DRB1 typing.
Because HLA-C is located between the A and B loci and because
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**Anti-HLA Antibody Response**

HLA antigens were originally identified with anti-

bodies induced by blood transfusion [14] or pregnancy
[15, 16]. Another source of immunization is transplan-
tation. For the formation of HLA antibodies, a confron-
tation with foreign HLA antigens is necessary, this is in
contrast to ABO antibodies that are naturally present. In
the past, when HLA antigens were identified serologi-
cally, the dogma was that a confrontation with foreign
HLA antigens will lead to the formation of antibodies,
although the exact epitopes recognized by the antibodies
were unknown. Molecular and high-resolution HLA
typing has led to the exact definition of the amino-acid
sequence of all HLA alleles and to the identification of
the important antibody epitopes on the different HLA
molecules [17]. These epitopes are not exclusive for one
particular HLA allele, but HLA alleles may share epi-
topes. As patients do not produce antibodies to epitopes
present on their own HLA antigens, one can even iden-
tify foreign HLA alleles that only carry epitopes shared
with the patient’s own HLA antigens and which will not
induce antibodies [18].
Anti-HLA Alloreactivity

Alloreactive T cells mediate graft rejection after solid organ transplantation (host anti-donor response), and graft-versus-host disease (donor anti-host response) in HSCT. Such alloresponses can be based on either a direct allorecognition of the incompatible HLA antigen through peptide-dependent structural mimicry [19], or an indirect allorecognition of donor HLA peptides presented by recipient HLA molecules [20]. The reactivity of T cells with incompatible HLA molecules is unexpected as T cells are educated to recognize viral and bacterial peptides in the context of self HLA. Recent studies have shown that reactivity with incompatible HLA is based on crossreactivity of T cells meant to recognize viral peptides in self HLA [21, 22]. As a consequence, a patient, who has never been confronted with foreign HLA but has experienced several infections, can have memory T cells to foreign HLA antigens on a graft affecting transplant outcome.

Impact of HLA Diversity on Organ Transplantation

Already in the early days of renal transplantation, it became clear that the HLA antigens are strong transplantation antigens [23]. The best transplant results were obtained when donor and recipient were HLA identical, and the more HLA mismatches, the poorer the results. The introduction of very potent immunosuppressive drugs has diminished the impact of HLA matching on graft survival, although the beneficial effect still exists [24]. A more important reason to match for HLA in solid organ transplantation is to prevent the formation of antibodies. The presence of donor-specific HLA antibodies before transplantation leads to a hyperacute rejection [25]. This complication is prevented by a serological crossmatch and the exclusion of donors towards whom the patient has formed specific antibodies [26]. However, HLA antibodies may also be formed after transplantation and are associated with acute and chronic graft rejection [27]. The more foreign antibody epitopes there are present on the donor organ, the higher the chance that a patient will form antibodies [28]. Especially for children, who often need a retransplantation, this is a potential problem as the presence of many HLA antibodies will prevent their retransplantation due to positive crossmatches [29]. A computer algorithm developed by Rene Duquesnoy, called HLAMatchmaker, can determine the immunogenicity of an HLA mismatch by comparing the epitopes on foreign HLA antigens with those on the patient’s own HLA antigens [30]. This tool is very useful for the selection of not very immunogenic HLA mismatches for children.

Highly sensitized patients, who have formed many HLA antibodies directed against many different HLA antigens, are very difficult to transplant as crossmatches with a potential donor are very often positive. For this reason these patients tend to accumulate on the transplantation waiting lists. A special strategy has been shown to be instrumental in facilitating transplantation of this difficult patient group [31]. It gives priority to these patients, when donors become available towards whom the patient do not produce antibodies, the so-called ‘acceptable mismatch program’ [31]. The HLAMatchmaker algorithm can help here, too, to define HLA antigens towards which a patient will not make antibodies, as illustrated in figure 3.

Impact of HLA Diversity on Donor Selection for HSCT

Allogenic HSCT is a well-recognized therapy for hematological malignancies and inborn errors of the immune system [32]. HLA polymorphism represents a ma-
major barrier to transplantation, because HLA incompatibilities at the allele level (down to a difference of one amino acid) can be efficiently recognized by alloreactive T lymphocytes. Therefore, the best HSC donor is a sibling who has inherited the same paternal and maternal HLA haplotypes as the patient. The probability to find a genotypically identical sibling donor is 25%, and it increases with the number of siblings: i.e. the probability is 43.7% with 2 siblings, 57.8% with 3 siblings (see fig. 2), or 68.4% with 4 siblings. However, because of a lack of matched sibling donors, at least in West-European populations, an increasing number of patients can have access to HSCT thanks to a worldwide registry of volunteer donors (BMDW: Bone Marrow Donor Worldwide). These donors are HLA typed, mostly at the low/intermediate-resolution level for HLA-A/B/(C)/DRB1/(DQB1), and they can be selected from the database on the basis of the best matching rate with a given patient. Searches are usually based on HLA allele compatibility for the HLA-A,B,C,DRB1,DQB1 loci, a so-called ‘10/10 match’ [33, 34]. Any mismatch results in a 9/10-matched donor, 2 mismatches in a 8/10-matched donor, etc. Because of the extensive HLA polymorphism, a 10/10-matched donor can only be identified for about 50% of all patients, and a 9/10-matched donor for an additional 20–30% of all patients [35–38] (table 1).

The implementation of recipient and donor HLA high-resolution genotyping in the clinical practice has clearly improved the success of transplantation through a better matching [33, 34, 39], but it has also resulted in more complex search algorithms. Several studies have shown that patients’ allele and haplotype frequencies have a strong impact on the chance to identify a compatible donor [37, 38, 40, 41]. Despite the fact that more than 20 million donors are now registered in BMDW, for about 5% of the patients there is no single potential donor in the database, either because the patient’s HLA-A,B,DRB1 phenotype is absent in the database or because available donors typed at the intermediate/high-resolution level have allele mismatches. As summarized in table 1, about 70% of all patients, mainly of European ancestry, may benefit from a 10–9/10-matched donor on the basis of high-resolution typing of potential donors selected from the international registry. The large registries with several million donors (USA, Germany) allow the identification of a compatible donor for a majority of patients of European ancestry. However, a number of patients rely on smaller registries to find a matched donor, because specific low-frequency HLA haplotypes present a more restricted geographical distribution (see below).

### Table 1. Patient/donor-matching probabilities in different European countries

<table>
<thead>
<tr>
<th>Patients with a 9–10/10-matched donor, %</th>
<th>Time frame, months&lt;sup&gt;d&lt;/sup&gt;</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>69%&lt;sup&gt;a&lt;/sup&gt;</td>
<td>2.5</td>
<td>[35]</td>
</tr>
<tr>
<td>72%</td>
<td>2.5–3.2</td>
<td>[42]</td>
</tr>
<tr>
<td>61.1%&lt;sup&gt;b&lt;/sup&gt;</td>
<td>&lt;1/1.5/16&lt;sup&gt;e&lt;/sup&gt;</td>
<td>[40]</td>
</tr>
<tr>
<td>78.3%&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1.84</td>
<td>[43]</td>
</tr>
<tr>
<td>81.8%</td>
<td>1.8/2.4/2.8&lt;sup&gt;e&lt;/sup&gt;</td>
<td>[37]</td>
</tr>
<tr>
<td>79%</td>
<td>n.a.</td>
<td>[38]</td>
</tr>
</tbody>
</table>

<sup>a</sup> <9/10 in 13% of all patients.  
<sup>b</sup> Corresponds to 10/10-matched donors only.  
<sup>c</sup> Exceptionally 8/10-matched donors.  
<sup>d</sup> Time from start of the search until the identification of a 9–10/10-matched donor.  
<sup>e</sup> Search duration for patients with high-/intermediate-/low-frequency HLA haplotypes.

### Table 2. Relative number of C<sup>*02:02</sup>-positive donors among 226 donors with the A<sup>*03:01</sup>-B<sup>*51:01</sup>-DRB1<sup>*04:01</sup> haplotype

<table>
<thead>
<tr>
<th>Registry</th>
<th>Donors&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Relative occurrence&lt;sup&gt;b&lt;/sup&gt;</th>
<th>C&lt;sup&gt;*02:02&lt;/sup&gt;-positive donors, n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Germany</td>
<td>90</td>
<td>1.84 × 10&lt;sup&gt;–5&lt;/sup&gt;</td>
<td>9 (10%)</td>
</tr>
<tr>
<td>USA</td>
<td>65</td>
<td>0.88 × 10&lt;sup&gt;–5&lt;/sup&gt;</td>
<td>4 (6%)</td>
</tr>
<tr>
<td>The UK</td>
<td>63</td>
<td>7.19 × 10&lt;sup&gt;–5&lt;/sup&gt;</td>
<td>0</td>
</tr>
<tr>
<td>The Netherlands</td>
<td>5</td>
<td>11.9 × 10&lt;sup&gt;–5&lt;/sup&gt;</td>
<td>1 (20%)</td>
</tr>
<tr>
<td>Portugal</td>
<td>3</td>
<td>0.94 × 10&lt;sup&gt;–5&lt;/sup&gt;</td>
<td>2 (66%)</td>
</tr>
</tbody>
</table>

<sup>a</sup> Number of donors with the phenotypes A3-B51-DR4/A1-B8-DR3, A3-B51-DR4/A3-B7-DR15 and A3-B51-DR4/A29-B44-DR7 with available HLA-C-typing results.  
<sup>b</sup> Relative occurrence of the 3 phenotypes A1/3,B8/51,DR3/4, A3/–,B7/51,DR15/4 and A3/29,B51/44,DR4/7 with available HLA-C types in each of the 5 national registries.

### HLA Diversity and Search Algorithm

What is the impact of HLA diversity in different European populations on donor searches? European stem cell donor registries have now more than 8 million HLA-typed donors, in the worldwide database of 21 million donors, distributed in 69 different national registries from 50 countries, characterized by their widely differing sizes (www.bmdw.org):

- registries with less than 100,000 donors: such as the Danish (28,000), Dutch (44,000), Swiss (48,000), or Austrian (63,000) registries,
Table 3. Relative frequencies of potential donors in two search categories

<table>
<thead>
<tr>
<th>Registry</th>
<th>1–15 donors</th>
<th>16–150 donors</th>
</tr>
</thead>
<tbody>
<tr>
<td>USA</td>
<td>29</td>
<td>24</td>
</tr>
<tr>
<td>Germany</td>
<td>19</td>
<td>23</td>
</tr>
<tr>
<td>Great Britain/Ireland</td>
<td>2</td>
<td>4.5</td>
</tr>
<tr>
<td>Brazil</td>
<td>19</td>
<td>20</td>
</tr>
<tr>
<td>Australia/Canada</td>
<td>2</td>
<td>3.5</td>
</tr>
<tr>
<td>Portugal/Spain/Italy</td>
<td>13</td>
<td>7</td>
</tr>
<tr>
<td>Belgium/France/Switzerland/the Netherlands</td>
<td>2</td>
<td>2.5</td>
</tr>
<tr>
<td>Sweden/Denmark</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Poland/Slovenia/Slovakia/Croatia</td>
<td>6</td>
<td>4.5</td>
</tr>
</tbody>
</table>

The results are the average ratio (in percent) of potential donors from any given national registry over the total number of potentially matched donors available for each of the 2 groups of patients (n = 168 for the searches with 1–15 donors, n = 2,594 for the searches with 16–150 donors).

All searches were run in 2012. The respective numbers of donors from other European, Central/South-American, and Asian registries are too small to be meaningful and are therefore not represented. See text for more details.

Bold values show higher relative frequencies of donors.

Discussion

Since the use of unrelated HSC donors is deemed to grow, the efficiency of a search algorithm will benefit from a larger diversity of HLA haplotypes represented in the database, and of a higher quality of the HLA-typing data of the newly registered donors. A few registries are now providing sequencing data at HLA-A,B,C,DRB1,DQB1 loci. A better knowledge of HLA allele/haplotype frequencies in different populations will certainly allow to better delineate search strategy and thus focus donor selection on specific donor population groups, whenever donors are not typed at a high-resolution level for all HLA loci. Particularly challenging are searches for patients of non-European ancestry because of the underrepresentation of donors from other ethnic groups in the BMDW registry. This problem is underlined by a recent study reporting a 33% success rate in donor identification (7–8/8 match) for patients of African origin. This observation was further supported by the analysis of the same A*03:01-B*51:01-C*02:02-DRB1*04:01 haplotype when combined with any of the most frequent haplotypes (A1-B8-DR3, A3-B7-DR15 and A29-B44-DR7). An interrogation of the BMDW registry showed that of 226 donors only 16 donors (7%) typed as B*51:01-C*02:02 (table 2). Again 100% of the 63 donors from the UK and 94% from the US registries were C-incompatible, whereas 2 out of 3 donors from the Portuguese registry were compatible for HLA-C (table 2).

In order to indirectly evaluate the impact of HLA diversity in different European populations, we determined the ratio of potential donors originating from ‘small’ or ‘intermediate-sized’ European registries compared to donors from large registries (USA, Germany, the UK). We thus compared the origin (national registry) of potential donors for the 2 following categories of patients: those patients with more common haplotypes, i.e. with more than 15 potential donors in the BMDW (40 searches), and those patients with less than 16 donors (28 searches). As shown in table 3 (numbers in bold), a trend towards a higher relative frequency of donors from South-European (Portugal, Spain, Italy), North-European (Sweden, Denmark) and East-European (Poland, Slovenia, Croatia, Slovakia) registries is observed. Interestingly, the relative percentage of BMDW-available donors from the British, Irish, German, Australian or Canadian registries was lower in searches for patients with low-frequency haplotypes.
ancestry, as compared to 76% for patients of European ancestry [44]. The increasing heterogeneity of the European populations, with patients of African, Asian, South-American, or mixed ancestry, calls for an increase in donor recruitment among these populations.

For those patients who cannot benefit from a fully matched donor, the challenge remains to identify more permissive or less immunogenic HLA mismatches, which are associated with less clinical complications. An alternative for these patients is HSCT from haploidentical donors or from cord blood units.

As a consequence of the extreme diversity of HLA-haplotypic associations in many populations, it appears of primary importance to support donor recruitment in all European countries. Even though the 2 largest registries account for more than 70% of all donors in Europe, they may not cover sufficient HLA diversity to provide matched donors, particularly for patients of South- and East-European ancestry. HLA diversity does not only mean the inclusion of new A,B,DRB1 phenotypes in the database, but also extending the allelic variability within already existing A,B,DRB1 phenotypes. The development of the so-called ‘small’ registries should be a priority, but should not be at the expenses of HLA-typing quality.

The situation is different in organ transplantation, where a worldwide search for HLA-compatible donors is not feasible for logistic reasons. Nevertheless, the best results here are also obtained with fully HLA-matched donors, although the outcome of HLA-mismatched transplants is still very good due to efficient immunosuppressive medication. The main reason for graft failure in renal transplantation is the (de novo) formation of donor-specific HLA antibodies [27].

If the induction of HLA antibodies can be prevented, the risk to develop acute or chronic graft rejection will diminish. Furthermore, the chance that a patient will receive a retransplantation depends on the degree of sensitization after the rejection of a previous transplant. Preventing the formation of HLA antibodies will be beneficial for patients in need for a retransplantation as is often the case in paediatric patients.

For these reasons, the strategy in organ transplantation will be more focussed on matching at the level of antibody epitopes [45]. Individual HLA alleles express different antibody epitopes, but these epitopes are shared with some other HLA alleles. This information together with the fact that a patient will not form antibodies to epitopes present on her/his own HLA molecules will be the basis of future matching strategies using algorithms like HLAMatchmaker. This will facilitate donor selection, as only 150–200 amino-acid variations within more than 5,000 HLA class I alleles are responsible for the induction of virtually all HLA antibodies. The current challenge is a clear definition of these crucial amino-acid variations associated with the induction of antibodies, and next the definition of the exact epitopes recognized by allo-antibodies in order to select the optimal donor for sensitized patients.

It has become clear in this review that the enormous polymorphism of HLA antigens, which is essential for our defence to viruses and bacteria, hampers the selection of an optimal donor for a patient in need for a transplantation. Knowledge of the exact amino-acid sequence of all HLA alleles, and especially the crucial positions for T cell and antibody recognition, and of the HLA-haplotype distribution in different populations all over the world will enable the selection of the best donor for a potential transplant recipient.

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References


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