A comparative reference study for the validation of HLA-matching algorithms in the search for allogeneic hematopoietic stem cell donors and cord blood units


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Key words
donor registry; hematopoietic stem cell transplantation; human leukocyte antigen; immunogenetics; matching algorithm; patient–donor matching

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Received 27 November 2015; revised 19 April 2016; accepted 22 April 2016
doi: 10.1111/tan.12817

Abstract

The accuracy of human leukocyte antigen (HLA)-matching algorithms is a prerequisite for the correct and efficient identification of optimal unrelated donors for patients requiring hematopoietic stem cell transplantation. The goal of this World Marrow Donor Association study was to validate established matching algorithms from different international donor registries by challenging them with simulated input data and subsequently comparing the output. This experiment addressed three specific aspects of HLA matching using different datasets for tasks of increasing complexity. The first two tasks targeted the traditional matching approach identifying discrepancies between patient and donor HLA genotypes by counting antigen and allele differences. Contemporary matching procedures predicting the probability for HLA identity using haplotype frequencies were addressed by the third task. In each task, the identified disparities between the results of the participating computer programs were analyzed, classified and quantified. This study led to a deep understanding of the algorithms participating and finally produced virtually identical results. The unresolved discrepancies total to less than 1%, 4% and 2% for the three tasks and are mostly because of individual decisions in the design of the programs. Based on these findings, reference results for the three input data sets were compiled that can be used to validate future matching algorithms and thus improve the quality of the global donor search process.

Introduction

Human leukocyte antigen (HLA) matching between patient and donor is the pivotal factor for the success of an allogeneic hematopoietic stem cell (HSC) transplantation (1–3). Therefore, the rapid and reliable identification of suitable adult volunteer HSC donor candidates and/or cord blood units for individual patients is of primary importance. This challenging search process is routinely performed in a donor registry or cord blood bank by a computer program in which the HLA-matching algorithm (HMA) can be regarded as the core element (4). For simplicity, we will use the term ‘donor’ to refer to donors of HSCs from bone marrow or peripheral blood and cord blood units and the term ‘donor registry’ shall include cord blood banks.

Historically, the HMAs were developed independently within each donor registry as a part of a highly specialized individual IT infrastructure (5). Subsequently, HMAs have been developed further to encompass the evolution of the serological and molecular HLA nomenclature and also because of the growing scientific understanding of the clinical matching requirements (6–9). In particular, the HLA typing resolution required, the number of HLA loci to be considered and the number of donors typed incompletely or at insufficient resolution have promoted a heterogeneous range of matching philosophies. The introduction of probabilistic concepts into the algorithms for predicting high-resolution HLA allele-level matching established the current state of the art in patient–donor matching (10–13).
International cooperation plays an essential role in the HSC donor search process. In 2014, 49% or 8112 of the 16,655 globally shipped bone marrow or peripheral blood stem cells products were imported (14). In this HSC exchange Bone Marrow Donors Worldwide (BMDW) allows for searching of the global repository of currently more than 26 million HSC donors using its own matching procedure (15). In addition, the European Marrow Donor Information System (EMDIS) is of major importance as its 35 member registries provide electronic access by their individual HMAs to 90% of the globally available HSC sources (16).

From the perspective of search coordinators, all individual HMA implementations should present a comparable picture of the donor situation for their patients. However, the long evolution of HMAs has not led to a convergence of their behavior, but instead an increasing number of differences between their algorithms produce diverse results, which can complicate the donor search process.

For this reason, the Matching Validation Subcommittee of the IT working group of the World Marrow Donor Association (WMDA) initiated two projects for the global validation and standardization of HMAs to improve the quality and consistency in global donor searches (17). The first was to define a formal specification of HLA matching using accurate terminology (18). The second effort presented here involves the comparison and cross-validation of matching algorithms by processing a panel of reference data sets and comparing the results obtained from the different registries’ algorithms. This approach had the potential to highlight so far uncharacterized differences between algorithms that cannot be detected based on a high-level description of their functional components and their underlying assumptions. The relevance of this unique comparative study is underlined by the fact that the largest HLA-matching providers worldwide have participated (see affiliations).

**Material and methods**

The Matching Validation Subcommittee defined three matching validation tasks (MVTs). For each task, a new matching validation data set (MVS) comprising the HLA typings of 1000 patients and 10,000 donors was created. The 10 million possible patient–donor pairs of these three MVSs had to be used by the participants as input data for their own HMA. The complexity of the computational tasks with regard to algorithmic requirements and considered data increased from MVT 1 to 3.

All materials and results necessary to run the experiments for newly developed or modified algorithms and to analyze and validate match results are made available for download as part of the Supporting Information. In the following, references to tables and figures in the Supporting Information are denoted with ‘S’.

**Simulation of ambiguous HLA genotypes**

The genotypes of the patients and the donors for the three tasks of this study were all separately generated by independently drawing two haplotypes from the set of high-resolution US Caucasian haplotypes with a probability corresponding to their frequencies given in Maiers et al. (19). For the purpose of having patient and donor genotypes at varying levels of typing resolution, the HLA used in the tasks corresponds to what the known HLA type would have been if those individuals had been typed using representative methods and typing kits. The ambiguities thus introduced are based on IPD-IMGT/HLA Database (20) releases v2.16.0 for MVS 1 and MVS 2 and v3.4.0 for MVS 3. Patient and donor HLA genotypes are encoded using genotype list (GL) string syntax (21) and multiple allele codes are used to reflect typing ambiguities (22). Identical amino acid sequences in the antigen recognition domain (ARD) were grouped together. The ARD groups have been defined on the basis of the two fields ‘g’-groups specifically introduced in the Common and Well-Documented (CWD) alleles catalog (23). The distribution of ambiguously or incompletely typed donors shapes a typical registry profile. Patients are usually typed at high-resolution upfront. However, in order to challenge the algorithms, a substantial amount of ambiguity has been introduced into the patient HLA assignments.

**Matching validation task 1 (mismatch counting)**

The goal of the first MVT was to compare the identification of definitive mismatches in ambiguous molecular HLA assignments. MVS 1 was generated based on 3-locus HLA-A~B~DRB1 haplotypes. Of the donor genotypes, 15% contained only allele assignments or alleles within a single ARD group, 24% had a combination of allele assignments and multiple allele codes and 61% contained first-field HLA typing results coded as XX-codes (24). Among the patient genotypes, the corresponding distribution was 49%, 39% and 12%. All donors and patients were typed for all three loci HLA-A, -B and -DRB1 (see Tables S1 and S2 for more details).

The task was to report for each patient–donor pair the total number of differences at each HLA locus. No distinction was to be made between antigen and allele mismatches and linkage disequilibrium was not to be considered.

**Matching validation task 2 (mismatch grading)**

The major refinement of the second task was the requirement to discriminate between mismatches at the antigen (serologic) level and those at the allele level in the counting. The distribution of the HLA assignments was similar to MVS 1 but, as a further complication, in MVS 2 serological assignments instead of XX-codes were used (see Table S1 for more details). The task was to report for each patient–donor pair at each HLA locus the total number of differences and the number of antigen differences. Again, linkage disequilibrium was not to be considered.
Matching validation task 3 (matching probability)

In addition to the assignment of matching classifications required in tasks 1 and 2, the third task required the calculation of allele-level match probabilities from ambiguously typed donor and patient HLA genotypes. Implementing such an HMA was a double challenge, first by virtue of the complexity of the task itself and second because of the requirement for computational runtime efficiency as the number of possible high-resolution haplotype pairs (diplotypes) can be extremely high for incomplete or insufficiently resolved typing data.

MVS 3 was generated based on the above mentioned US Caucasoid 5-locus HLA-A~C~B~DRB1~DQB1 haplotypes using a mixture of HLA typing methods that are typically found in registries with many years of accumulated HLA typings. Here, 6% of the donor genotypes contained only allele or ARD group assignments, 33% had a combination of allele assignments and multiple allele codes and 61% contained only XX-codes. Among the patient genotypes the corresponding distribution was 71%, 15% and 14%. All patients were typed for all five HLA loci, however, typing for the loci HLA-C, -DRB1 and -DQB1 was removed for 80%, 10% and 90% of the donors, respectively, to more closely resemble the practical situation and-DQB1 was removed for 80%, 10% and 90% of the donors, all five HLA loci, however, typing for the loci HLA-C, -DRB1 and -DQB1 was removed for 80%, 10% and 90% of the donors, respectively, to more closely resemble the practical situation occurring in most donor registries (see Table S1). By design, the genotypes of all individuals were explainable by the haplotypes contained in the frequency table provided.

On the basis of the same 5-locus haplotype frequencies used for the simulation of the patient and donor population for each patient–donor pair the overall 9/10 and 10/10 matching probability as well as the locus-specific 2/2 matching predictions each accompanied by a match grade character had to be computed.

General requirements

The resulting file for each MVT had to contain the specific data items for all 10^7 possible patient–donor pairs in a specific format (see Figure 1). The early stages of the analysis of MVT 1 and 2 showed that very strict and precise specification of the task was necessary to get comparable files. For this purpose, we defined that alleles within the same ARD group had to be considered an allele match. The counting of mismatches for HLA molecular assignments had to be implemented according to the #Max column of Table S3, which was derived from the WMDA HLA matching framework (18).

For MVT 2 a common mapping of alleles to antigens and vice versa was important for obtaining comparable results. For this purpose, the DNA-to-serology mappings defined in the WMDA file rel_dna_ser.txt (25) according to IPD-IMGT/HLA Database release v3.6.0 have been used. To ensure consistency with World Health Organization (WHO)-assigned antigen mappings it was decided to disregard the expert-assigned values of the WMDA mapping file in this experiment.

The differences observed in a first round of analysis for MVT 3 showed the need for an even stricter rule set to achieve comparable results. In summary, the requirements to achieve a ‘baseline’ MVT 3 result are:

R1 ARD level matching has to be used.
R2 Mismatches have to be counted according to Table S3.
R3 WHO HLA nomenclature with WMDA extensions has to be used (24).
R4 The reported matching probabilities have to utilize simple rounding half up to integers and range between 0 and 100.
R5 Possible values for the locus-specific match grade characters are A (allele-level match), P (potential allele-level match) and M (mismatch).
R6 When a locus is not typed, the matching probability and the match grade character have to be calculated and provided.
R7 The probability for a 9/10 match refers to exactly the probability for a 9/10 match, not for a 9/10 match or better.
R8 The locus-specific results have to be calculated on the basis of the possible diplotypes, i.e. linkage disequilibrium has to be considered.
R9 The locus-specific results must differentiate between match probabilities that are exact values and those that are the result of rounding. Full allele-level matches (A) that lack mismatching alternative genotypes are given a result of ‘A;100’. Potential allele-level matches (P) that do have mismatching genotypes of low likelihood, such that the match probability rounds up to 100, are given a result of ‘P;100’. Likewise, complete mismatches (M) with no possible overlapping genotypes are given a result of ‘M;0’. However, a result of ‘P;0’ is given to cases where overlapping genotypes do exist and the match probability rounds down to 0.
R10 The calculation of the matching predictions has to consider all theoretically possible haplotypes/diplotypes (i.e. no trimming of the set of possible diplotypes).
R11 Multiple allele codes that have been discontinued (26) have to be reasonably reinterpreted within the current HLA nomenclature.

Comparison and statistics

In the central analyses for MVT 1 and 2, the overall consistency of the data provided and their compliance with the specification was initially checked. Then, for each patient–donor pair the reported numbers of match differences from the submitted result files were compared separately by means of Perl (27) scripts in which the redundant cases per locus were only considered once. In case of disparity, the values provided by the participants were collated in dedicated spreadsheets in which the cells containing values deviating from the majority (if any) were displayed in color. The discrepancies were manually inspected and categorized. These preliminary findings were discussed and, when necessary, clarified and adjusted in the meetings of the Matching Validation Subcommittee in order to compile a consensus result.

For MVT 3, a two-step analysis approach was used. First, an overview comparison between the results of every pair of
The total number of HLA differences (allele or antigen) for the loci HLA-A, -B and -DRB1 had to be reported (3 values). The valid range for the differences is 0 to 2.

The total number of differences (allele or antigen) for the loci HLA-A, -B and -DRB1 as well as the number of antigen differences had to be reported (6 values). The valid range for the differences is 0 to 2. For each locus the total number of differences must be greater or equal to the number of antigen differences.

A match grade character (A=allele match, P=potential allele match, M=mismatch) and the 2/2 match probability for the loci HLA-A, -C, -B, -DRB1 and -DQB1, as well as the overall 9/10 and 10/10 match probability had to be reported (12 values).

Figure 1 Result file formats for matching validation task (MVT) 1–3 (A to C). A single line had to be reported for each of the $10^7$ possible patient–donor pairs using semicolon as field separator.

Participants was made in which only the number of discrepant results concerning the three areas ‘overall 10/10 and 9/10 matching predictions’, ‘locus-specific 2/2 matching predictions’ and ‘locus-specific match grade characters’ were counted. Second, for the detailed analysis Perl scripts were used to collect the specific result items for all discrepant cases in suitably structured spreadsheets including the observed differences and the number of possible diplotypes as indicator for the complexity of the considered patient–donor pair. If necessary, differences concerning probability values were studied in detail by a trace program dissecting the genotypes of a patient–donor pair into their diplotypes and their individual probabilities. This allowed to analyze and classify all occurring differences in detail before visualizing them using the R statistics software package.

Characteristics of the HMAs compared

Some characteristics of the matching programs contributed by the participating groups are summarized in Table S4. These
Table 1 Observed disparities between the seven results submitted for MVT 1

<table>
<thead>
<tr>
<th></th>
<th>Total</th>
<th>B</th>
<th>DRB1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unique HLA typing pairs</td>
<td>123,708</td>
<td>580,659</td>
<td>229,504</td>
</tr>
<tr>
<td>Discrepant cases between all result files</td>
<td>857</td>
<td>2450</td>
<td>1951</td>
</tr>
<tr>
<td>Discrepant cases between all result files (%)</td>
<td>0.69</td>
<td>0.42</td>
<td>0.85</td>
</tr>
</tbody>
</table>

*The number of unique locus-wise typing pairs, the number of discrepant cases and the percentage of discrepant cases are shown for the total number of differences (tot) for the loci HLA-A, -B and -DRB1.

results and the percentage of discrepant results are shown for the total number of results to unique HLA typing pairs for each locus. The showsthenumberofdisparitiesafterreducingthetotalnumberhavethesameoutcomevariablesin each result file. Table 1 and the 10,000 donors were identified and combined into unique for each locus repeating HLA types within the 1000 patients to control the results for consistency and to facilitate the comparison of discrepancies for the number of antigen mismatches was observed across all participants in identifying allele mismatches based on serological assignments. The strikingly higher numbers of discrepancies for the number of antigen mismatches was caused by several different approaches in the treatment of serologic assignments. Overall, the differences observed could be attributed to the following reasons:

1. Usage of different DNA-to-serology mapping tables. This is actually a violation of the specification but some participating algorithms could not be adapted to use a configurable mapping table with reasonable effort and time (see also Table S3).
2. Different decisions on potential allele matches between a serological and a molecular assignment. Those problems occur when broad and split/associated antigens are mixed in the serology-DNA-correspondence table entries considered, e.g. donor A25 which is a split antigen of A10 vs patient A*26:ADZV, containing A*26:15 (A10).
3. Different decisions on potential antigen matches between two molecular assignments. Those problems also occur when broad and split/associated antigens are mixed in the serology-DNA-correspondence table entries considered, e.g. donor A*25:01 (A25) vs patient A*26:ADZV, containing A*26:15 (A10).
4. Different treatment of apparent serological mismatches in the context of alleles bridging serological families, e.g. a donor A24 is an apparent antigen difference with patient A3, however because A*24:18 is showing A24/A3 as corresponding serology, this pair could be classified as a potential allele match. As a consequence, those cases were even classified as potential antigen matches by HMA #5.
5. Implementation specific features, e.g. no differentiation between B64/65, DR13/14 and DR15/16, i.e. those split antigens are generally mapped back to their broad value. In other words, apparent serologic split differences for such cases were not reported as antigen mismatches.

Table 1 illustrates the number and kind of the remaining disparities after reducing the total number of results to unique pairs of HLA types for each locus.

Surprisingly, the disparities for the total number of differences were slightly lower than in MVT 1. This can be explained by algorithmic improvements of the implementation discrepancy in task 1 that are overcompensating some new discrepancies among all participants in identifying allele mismatches based on serological assignments. The strikingly higher numbers of discrepancies for the number of antigen mismatches was caused by several different approaches in the treatment of serologic assignments. Overall, the differences observed could be attributed to the following reasons:

1. Usage of different DNA-to-serology mapping tables. This is actually a violation of the specification but some participating algorithms could not be adapted to use a configurable mapping table with reasonable effort and time (see also Table S3).
2. Different decisions on potential allele matches between a serological and a molecular assignment. Those problems occur when broad and split/associated antigens are mixed in the serology-DNA-correspondence table entries considered, e.g. donor A25 which is a split antigen of A10 vs patient A*26:ADZV, containing A*26:15 (A10).
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4. Different treatment of apparent serological mismatches in the context of alleles bridging serological families, e.g. a donor A24 is an apparent antigen difference with patient A3, however because A*24:18 is showing A24/A3 as corresponding serology, this pair could be classified as a potential allele match. As a consequence, those cases were even classified as potential antigen matches by HMA #5.
5. Implementation specific features, e.g. no differentiation between B64/65, DR13/14 and DR15/16, i.e. those split antigens are generally mapped back to their broad value. In other words, apparent serologic split differences for such cases were not reported as antigen mismatches.

Although the rate of concordance achieved was quite high, for the above reasons the compilation of a true consensus file

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Table 2 Observed disparities between the six results submitted for MVT 2

<table>
<thead>
<tr>
<th></th>
<th>A tot</th>
<th>A ag</th>
<th>B tot</th>
<th>B ag</th>
<th>DRB1 tot</th>
<th>DRB1 ag</th>
</tr>
</thead>
<tbody>
<tr>
<td>Unique HLA typing pairs</td>
<td>124,950</td>
<td>124,950</td>
<td>592,800</td>
<td>592,800</td>
<td>232,311</td>
<td>232,311</td>
</tr>
<tr>
<td>Discrepant cases between all result files</td>
<td>661</td>
<td>4774</td>
<td>2751</td>
<td>14,358</td>
<td>1018</td>
<td>5573</td>
</tr>
<tr>
<td>Discrepant cases between all result files (%)</td>
<td>0.53</td>
<td>3.82</td>
<td>0.46</td>
<td>2.42</td>
<td>0.44</td>
<td>2.40</td>
</tr>
</tbody>
</table>

aThe number of unique locus-wise typing pairs, the number of discrepant cases and the percentage of discrepant cases are shown for the total number of differences (tot) and the number of antigen differences (ag) for the loci HLA-A, -B and -DRB1.

HLA, human leukocyte antigen; MVT, matching validation task.

Table 3 Number and percentage (in brackets) of discrepant patient–donor pairs (1 × 10^7 data items) for any two participants concerning the locus-specific match grade characters (A/P/M) for all five HLA loci for MVT 3

<table>
<thead>
<tr>
<th>#1</th>
<th>#2</th>
<th>#3</th>
<th>#4</th>
<th>#5</th>
</tr>
</thead>
<tbody>
<tr>
<td>#1</td>
<td>×</td>
<td>9,356,595</td>
<td>3681</td>
<td>9,356,505</td>
</tr>
<tr>
<td>(93.6)</td>
<td>(&lt;0.1)</td>
<td>(93.6)</td>
<td>(&lt;0.1)</td>
<td></td>
</tr>
<tr>
<td>#2</td>
<td>×</td>
<td>9,355,983</td>
<td>52</td>
<td>9,356,505</td>
</tr>
<tr>
<td>(93.6)</td>
<td>(&lt;0.1)</td>
<td>(93.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>#3</td>
<td>×</td>
<td>9,355,983</td>
<td>3677</td>
<td>9,356,505</td>
</tr>
<tr>
<td>(93.6)</td>
<td>(&lt;0.1)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#4</td>
<td>×</td>
<td>9,356,505</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(93.6)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#5</td>
<td>×</td>
<td></td>
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</table>

HLA, human leukocyte antigen; MVT, matching validation task.

Results of MVT 3

Participants #1 to #5 provided results for the third MVT. The format of the result file is illustrated in Figure 1C. Overview comparisons of the match grades, overall match probabilities and locus-specific match probabilities of the participating HMAs are presented here. To support the validation of new HMA implementations, we also provide a more detailed comparative analysis with HMA trace output in Appendices S1 and S2, along with a consensus result file in Material S3.

Overview comparison of match grade characters

The comparison of the locus-specific match grade characters shows virtually identical results for participants #2 and #4 and for participants #1, #3 and #5, respectively (see Table 3). The large discrepancy observed was caused by the latter group not complying with requirement R9, i.e. did not distinguish between rounded and exact values for 0% and 100%. The 52 discrepant cases found within the first group could be tracked down to the different treatment of discontinued multiple allele codes (compliance with R11).

Overview comparison of match probabilities

The findings of the comparison of the overall and locus-specific probability values are shown in Tables 4 and 5. The subsequent detailed analysis of the disparities between each pair of participants allowed assigning them to the following reasons corresponding to the superscripts used in these tables:

1 Discrepant mismatch counting between a homozygous patient and a donor with a null allele or vice versa cases (cf. AA – AN in Table S3).
2 Discrepant mismatch counting when patient and donor are homozygous (cf. AA – BB in Table S3).
3 Trimming of the set of possible diplotypes.
4 Treatment of discontinued multiple allele codes (cf. R10).
5 Numerical artifacts due to floating point arithmetic in combination with rounding.
6 Conditional locus-specific probabilities.

Overall probabilities

The design of MVT 3 yielded a large number of patient–donor pairs with more than one mismatch implying 9/10 and 10/10 probabilities of zero. For the remaining, almost 14,000 pairs an excellent rate of concordance was achieved (see Table 4). In particular, the results of algorithms #2 and #4 were completely identical and the results of algorithms #1 and #3 were almost identical. The detailed analysis revealed that reason 1 and/or 2 above were the main cause of the higher deviations observed (see Figure 2). The slightly higher rate of disparities for participant #5 could be traced back to reason 3 above. This algorithm was impossible and the result file provided for reference in the Supporting Information reflects the discrepancies observed.
Table 5 Number and percentage (in brackets) of discrepant patient–donor pairs (1 × 10^7 data items) for any two participants concerning the 2/2 locus-specific predictions for all five HLA loci for MVT 3

<table>
<thead>
<tr>
<th></th>
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<th>#3</th>
<th>#4</th>
<th>#5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>× 729 1,2,5</td>
<td>n/a 6 (&lt;0.1)</td>
<td>678 1,2,5</td>
<td>186,919 2,3,5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#2</td>
<td>× n/a 6</td>
<td></td>
<td>225 4,5</td>
<td>186,453 3,5</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#3</td>
<td>× n/a 6</td>
<td>× n/a 6</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#4</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>#5</td>
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</table>

HLA, human leukocyte antigen; MVT, matching validation task; n/a, not applicable.

Superscripts indicate reasons for discrepancies as: 1, discrepant mismatch counting between a homozygous patient and a donor with a null allele or vice versa cases; 2, discrepant mismatch counting when patient and donor are homozygous; 3, trimming of the set of possible diplotypes; 4, treatment of discontinued multiple allele codes; 5, numerical artifacts due to floating point arithmetic in combination with rounding; 6, conditional locus-specific probabilities.

had to maintain a trimming threshold to deal with the computational complexity in cases with a high number of possible diplotypes. Actually, the output of algorithm #5 is fully identical to the results of algorithms #2 and #4 when excluding the 1005 donors without HLA-DRB1 assignments.

Locus-specific probabilities

For the probability of locus identity, only a few easily explainable disparities were found (see Table 5). Here, algorithms #2 and #4 altogether showed 225 cases with a deviation of one percentage point in the detailed analysis. Thirteen of these disparities were caused by reason 3 and the rest by reason 5 mentioned above. This is also true for most of the disparities with algorithm #1. However, the discrepant counting addressed in reasons 1 and 2 led to some higher deviations (see Figure 3). The substantial higher rate of disparities observed for participant #5 is again mainly caused by reason 3. When excluding the 1005 donors without HLA-DRB1 data, the result of algorithm #5 becomes virtually identical to #1, #2 and #4. The locus-specific probability values provided by participant #3 were not comparable to others because this algorithm returns probabilities for potentially 9/10 matched donors that are conditional on being exactly one mismatch.

Consensus result

In the course of the repeated comparison of gradually refined and/or corrected result files, it finally became apparent that the outputs of all algorithms were converging mainly because the
participants started to reduce or altogether avoid the trimming of long diplotype lists. Adapting to the other baseline require-
ments turned out to be too laborious for some implementations, especially because the advantage for daily use would only be marginal. Since algorithm #2 observed all baseline require-
ments, it was chosen as reference for the detailed analysis process. This way, all disparities could be explained satisfactorily and have been assessed by the members of the Matching Val-
ification Subcommittee. The result of participant #2 is provided
as consensus file for MVT 3 in the Supporting Information.
With regard to the accuracy of matching predictions provided
in the consensus file, the group agreed that the observed intrin-
sic deviations of 1% caused by floating point arithmetic (29) in combination with rounding are negligible.

Discussion

The objective of the project was to compare key aspects of
the behavior of HMAs when they have to deal with the wide
variety of HLA genotype data in today’s donor registries and
to use the results to identify relevant problems and pitfalls in
this complex task. The ultimate goal was to reach a consen-
sus on important underlying principles and the desirable results
wherever possible and to otherwise identify the key design deci-
sions where the algorithms or implementations may deliber-
ately deviate from each other. In those cases, the exercise served
as a cross-validation to ensure that the discrepancies between
the HMAs are restricted to the intended effects. While HSC
donor selection is a complex process that also must consider
many factors outside of HLA matching (30), we focused solely
on the contribution of HLA match calculations by HMAs to this
process.

Other methods of software validation are either impractical
(formal methods, code inspection) or less meaningful (exami-
nation of specification and documentation) than the functional
testing based on a large range of simulated practical input sig-
nals carried out in this study. In contrast to HLA typing metho-
donology, there are no established regulatory frameworks in place
for HMAs. However, previous validation efforts for the esti-
mation of HLA haplotype frequencies by the WMDA ITWG
Registry Diversity working group (31) are not only a building
block for this study but are also used in the validation of HLA
typing techniques (32).

The major challenges were related to the handling of the
complexity of HLA nomenclature in its historical context and
to the dealing with complexity of the haplotype-related cal-
culations for predictive matching. The sequence of the three
tasks was necessary to isolate groups of individual problems
from each other and to acquire an increasing understand-
ing of the individual design decisions underlying different
behaviors.

The first two MVTs addressed the locus-wise matching
decisions between patient and donor without considering fre-
quency information or linkage disequilibrium. In other words,
the algorithms were supposed to evaluate potential matches
using purely combinatorial methods. Strictly followed, such an
exhaustive approach leads to unusable search reports due to
extreme unlikely pairs like \textit{HLA-A*02:XX} potentially matching
\textit{HLA-A*03:XX} since both generic groups comprise null alleles
(case AN – AN in Table S3 is a match). Apparently, all partic-
ipating algorithms are intrinsically using heuristics most likely
based on allele frequencies or a set of CWD alleles which all
lead to identical results for MVT 1.

For MVT 2, however, the differences in the algorithms
became apparent since the correspondence between sero-
logical and molecular assignments is not unequivocal and well
defined for all alleles: some alleles have no official sero-
logical correspondence, some have several and some are only
attributed to a broad serological specificity. The combinatorial
consequences for matching between two serological assign-
ments or between a serological and molecular assignment as
well as for assigning a molecular mismatch to the antigen
or allele level are quite complex and have been described in
details in the result section. Apparently, cases like the examples
shown there cannot be decided satisfactorily based on the
currently available reference tables (25). As a consequence,
MVT 2 did not lead to a proper consensus and the result
summary documents a certain degree of variability. MVTs 1
and 2 reflect the dilemma between the historically developed
HLA nomenclature that is still the basis of current paradigms
in donor selection and the practical requirements in clinical
decision-making.

MVT 3 introduced another level of complexity by requiring
the use of haplotype frequencies to distinguish between likely
and unlikely potential matches. In the early rounds of analy-
sis, the large number of various discrepancies observed was
caused by incomplete or partially disregarded baseline require-
ments. Later stages showed that adhering to those rules and
assumptions makes all relevant discrepancies disappear and
lead to a consistent consensus result for MVT 3. This conver-
gence of results required modifying the configuration or imple-
mentation details of several participating matching algorithms
according to the requirements R1 – R11. As a consequence, this
exercise does not necessarily reflect their behavior in daily rou-
tine where for certain reasons other preferences may be given
priority.

In particular, the trimming of the set of possible diplotypes
is a usual approach to reduce the computational complexity
to save a substantial amount of processing time and give the
user an improved response time for display of a new search
report. Although in most cases this still gives reasonably pre-
cise results, it became apparent that it is impossible to give
estimates for the maximum error introduced. This error in
the matching probability returned can actually be quite sig-
nificant and therefore any trimming threshold must be chosen
with great care (see Appendix S1). Still, the improvement of
the performance of the HMAs will remain a major topic in
the refinement of all existing matching programs as long as
incomplete and ambiguous donor types have to be dealt with. Their efficiency not only has a major impact on the perceived responsiveness of user interfaces but also for the daily automatic update of thousands of search reports required for highly automated registry services or in the EMDIS communication system (33, 34). Deployment of high-performance computing clusters and development of HMAs that use parallel computing paradigms may allow registries to meet these big data challenges.

The participants have benefitted in multiple ways from this series of experiments. First, several errors became apparent and were fixed, but moreover many other more or less intended properties underwent a critical review. So eventually the quality of all HMAs was confirmed in certain aspects and improved in others. Second, future updates of all HMAs can refer back to the consensus results to ensure that no regressions or unintended features have been introduced. Similarly, the developers of new HMAs can validate their implementation using the Supporting Information accompanying this article. Lastly, the baseline result of MVT 3 allows for the measurement of the impact of any performance tweaks used in the real-life version of an HMA on the speed and quality of the results to make a sound cost–benefit judgment. Eventually, it will be up to the registry community cooperating within the WMDA to decide to which extent the MVTs can become a part of their global assurance efforts (35). Moreover, the experience gained in this study could be incorporated into the specifications for the HMA of the global search system of BMDW.

This study provides the first major contribution to the practical validation of HMAs, but it still leaves certain aspects uncovered and will have to be refined when HMAs further evolve. The most relevant limitation of this study is the fact that all patient and donor genotypes are simulated from the same set of haplotypes that are later used with a positive frequency. However in practice, a substantial fraction of the patients and more often the donors cannot be explained by the set(s) of haplotypes with known positive frequency. All probabilistic HMAs need fallback strategies for such situations that were not addressed in this experiment.

More challenges will arise when haplotype frequency tables specific to a population or a population subgroup become available and individual patients or donors cannot be clearly assigned. Such an approach would model the reality of matching in a global donor pool more realistically (36), but the ultimate benchmark for all haplotype frequency estimation efforts is the validation of real-matching predictions with typing outcome data. Nevertheless, the findings of this study are independent from the population haplotype frequencies used since the causes of the differences observed are either algorithmic or intrinsic to the HLA nomenclature.

Although such comparative analyses do not prove the correctness of any program, they do provide a strong indication of correctness due to the consensus results of independent implementations. The authors are aware that ‘even when the experts all agree, they may well be mistaken’ and that ‘when the experts are agreed, the opposite opinion cannot be held to be certain’ (37).

Acknowledgments

LG, JS and MM have been funded by Office of Naval Research Grant N00014-13-1-0039.

Conflicts of interest

The authors have declared no conflicting interests.

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Supporting Information

The following supporting information is available for this article:

Table S1. Characteristics of the matching validation tasks and associated data sets.
Table S2. Types of HLA assignments used and their resolution.
Table S3. Counting mismatches for 2 by 2 comparisons of patient and donor molecular HLA assignments.
Table S4. Software and hardware environments of the HMA implementations.
Appendix S1. PDF file with the detailed analysis of MVT 3 (including Figures S1–S3).
Appendix S2. Tar GZip file with an example trace for MVT 3 as Excel file.
Material S1. Tar GZip file containing README, MVS 1 and consensus result.
Material S2. Tar GZip file containing README (including the coding of disparities), MVS 2, DNA-to-serology mapping table and reference result.
Material S3. Tar GZip file containing README, MVS 3, list of ARD groups, haplotype frequency table and consensus result.