



HHS Public Access

Author manuscript

Hum Immunol. Author manuscript; available in PMC 2019 February 01.

Published in final edited form as:

Hum Immunol. 2018 February ; 79(2): 77–86. doi:10.1016/j.humimm.2017.12.004.

Collection and Storage of HLA NGS Genotyping Data for the 17th International HLA and Immunogenetics Workshop

Chia-Jung Chang¹, Kazutoyo Osoegawa², Robert P. Milius³, Martin Maiers³, Wenzhong Xiao^{1,4}, Marcelo Fernandez-Vi a^{2,5}, and Steven J. Mack⁶

¹Stanford Genome Technology Center, Palo Alto, CA, USA

²Histocompatibility, Immunogenetics & Disease Profiling Laboratory, Stanford Blood Center, Palo Alto, CA, USA

³Bioinformatics Research, National Marrow Donor Program, Minneapolis, MN, USA

⁴Massachusetts General Hospital and Shriners Hospital for Children, Boston, MA, USA

⁵Department of Pathology, Stanford University Medical Center, Stanford, CA, USA

⁶Center for Genetics, Children's Hospital Oakland Research Institute, Oakland, CA, USA

Abstract

For over 50 years, the International HLA and Immunogenetics Workshops (IHIW) have advanced the fields of histocompatibility and immunogenetics (H&I) via community sharing of technology,

Corresponding Author: Steven J. Mack, Center for Genetics, Children's Hospital Oakland Research Institute, Oakland, CA 94609, USA, sjmack@chori.org.

ONLINE RESOURCES

- A. BioSharing.org record for International HLA and Immunogenetics Workshop XML; <https://biosharing.org/bsg-s0007001> Accessed April 24, 2017.
- B. Instructions for using the 17th IHIWS Database; https://ihiws17.stanford.edu/ihiw_docs/17WS_Database_Manual_Registration_v1.pdf; Accessed April 25, 2017.
- C. Login portal for the 17th IHIW Database; <http://workshop.ihiws.org/>; Accessed April 25, 2017.
- D. Instructions for entering & uploading IHIW Typing reports; http://ihiws.org/wpcontent/uploads/2017/02/Instructions-for-entering_uploading-IHIWS-Typing-report_Version7.pdf; Accessed April 25, 2017.
- E. NMDP/Be The Match Bioinformatics Research GitHub repository; <https://github.com/nmdp-bioinformatics>; Accessed April 25, 2017.
- F. 17th IHIW GitHub repository; https://github.com/IHIW/bioinformatics/tree/master/db_related; Accessed December 10, 2017; referenced files are in the “/data”, “/hlaPoly”, and “/scripts” directories.
- G. IPD-IMGT/HLA Database FTP site; <ftp://ftp.ebi.ac.uk/pub/databases/ipd/imgt/hla/>; Accessed April 25, 2017; some referenced files are in the “/wmda” directory.
- H. hlaPoly: Identifying Novel Polymorphisms in HLA Sequences; <http://hlapoly.immunogenetics.org>; Accessed April 25, 2017.
- I. PyPop GitHub repository; <https://github.com/alexlancaster/pypop>; Accessed May 28, 2017.
- J. Feature service web-interface; <http://feature.b12x.org>; Accessed May 15, 2017; the Feature service allows individual gene feature sequences to be registered, returning an accession number for that sequence, and dereferences accession numbers to identify specific gene feature sequences; code is available on the NMDP GitHub repository under “service-feature”.
- K. GFE service; <http://gfe.b12x.org>; Accessed May 15, 2017; the GFE service accepts full-gene or multi-feature consensus sequence, splits it into individual features, which are registered with the Feature service, and returns a GFE notation; code is available on the NMDP GitHub repository under “service-gfe- submission”.
- L. GFE Allele-Calling Tool; <http://act.b12x.org/>; Accessed May 18, 2017; the Allele-Calling Tool accepts full-gene consensus sequence, and identifies the closest matching HLA allele and corresponding GFE notation; code is available on the NMDP GitHub repository under “service-act”.

Publisher's Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

experience and reagents, and the establishment of ongoing collaborative projects. Held in the fall of 2017, the 17th IHIW focused on the application of next generation sequencing (NGS) technologies for clinical and research goals in the H&I fields. NGS technologies have the potential to allow dramatic insights and advances in these fields, but the scope and sheer quantity of data associated with NGS raise challenges for their analysis, collection, exchange and storage. The 17th IHIW adopted a centralized approach to these issues, and we developed the tools, services and systems to create an effective system for capturing and managing these NGS data. We worked with NGS platform and software developers to define a set of distinct but equivalent NGS typing reports that record NGS data in a uniform fashion. The 17th IHIW database applied our standards, tools and services to collect, validate and store those structured, multi-platform data in an automated fashion. We have created community resources to enable exploration of the vast store of curated sequence and allele-name data in the IPD-IMGT/HLA Database, with the goal of creating a long-term community resource that integrates these curated data with new NGS sequence and polymorphism data, for advanced analyses and applications.

Keywords

International Workshop; 17th IHIW; Next Generation Sequencing; HLA; Database; Data Management; XML; HML

1. Introduction

1.1. The Histocompatibility Workshops

Since their introduction in 1964, the Histocompatibility Workshops have been forums for the exchange of community knowledge and experience, allowing histocompatibility and immunogenetics (H&I) researchers, clinicians and technologists to evaluate new methods and technologies, establish standards and advance ongoing collaborative projects. Sixteen International HLA and Immunogenetics Workshop (IHIW) meetings have been held on five continents over the last half-century[1–16], and the 17th IHIW was held in northern California in the fall of 2017, continuing many long-standing workshop projects.

The advent of next-generation sequencing (NGS) based genotyping technologies has allowed new insights and innovations for the fields of histocompatibility, immunogenetics and immunogenomics. The 17th IHIW's ultimate goals were to advance H&I basic research and clinical efforts through the application and evaluation of NGS HLA and KIR genotyping technologies, and to foster the development of NGS technologies tailored to meet the H&I community's needs, building on the technological and scientific momentum of the previous sixteen workshops.

Toward those ends, we developed systems, standards and tools for the collection, storage and management of NGS HLA genotyping data (the HLA genotype and associated consensus sequences) generated for 17th IHIW projects. The goals of this effort were to build on the data-collection and -storage experiences of previous workshops, and produce NGS data-managing tools that will support IHIW efforts and persist as public resources after the 17th IHIW. Here, we provide a brief overview of the challenges faced in organizing coordinated

data-generation and -collection efforts, the strategies we applied, and the tools, standards and services we developed to address these challenges.

1.2. The Challenges of Coordinated Data Collection

The collection, storage and analysis of data have been key issues of all workshops. Many workshops have used centralized databases[17–21], while in several of the more recent workshops, individual components and projects were responsible for collecting, managing and analyzing data[22–32]. Centralized data-management requires close communication between workshop participants and leaders, instrument and software vendors, and database developers to achieve consensus regarding required data content, data formats, reporting guidelines and quality standards. Sufficient time is also required for all parties involved to develop both the systems and tools to manage data, and the preliminary data on which to test the tools.

1.2.1. Reference Data Management—The specifics of the H&I field bring additional challenges that any data-management and analysis approach, centralized or decentralized, must address[33]. The body of HLA sequence data and associated allele names curated by the IPD-IMGT/HLA Database[34] (Reference Database) increases every four months; because workshop data-generation efforts often span multiple years, the details of the pertinent Reference Database version under which each HLA genotype was generated must be collected along with the genotyping data. The collection and management of such genotyping meta-data (Table 1) can be just as important for the workshop effort as the genotyping data themselves; without them it may not be possible to determine the extent to which datasets generated years apart or using different methods are equivalent. When workshop efforts span time periods that include major changes to the nomenclature[35, 36], these problems are only compounded.

1.2.2. Primary Data Management—The nature of the primary or “raw” data, from which all experimental data and meta-data are ultimately derived, can vary widely from method to method and from project to project. This was particularly pronounced for the molecular genotyping methods applied in the 11th through the 16th workshops, where multiple reference strand conformation analysis (RSCA), sequence-specific (SS) oligo (SSO), reverse SSO (rSSO), SS priming (SSP) and sequence-based typing (SBT) methods were in use, each with its own distinct type of primary data.

1.2.3. Allele Name Data Management—Allele name data must be recorded and managed in a standard manner to facilitate meaningful data-analysis. For many previous workshops, the management of HLA allele names was performed by humans, and involved data recorded in paper documents or spreadsheets in a variety of different ways. Humans are adept at “figuring out” the true meaning of unusual notations and spreadsheet-initiated errors that may occur, but machines are not. For example, “HLA-A*02:99” and “HLA-A*03:01:02” are often recorded as “02:99” or “03:01:02” in spreadsheet columns labeled “HLA-A”, “A”, etc.; however, common spreadsheet applications may change “02:99” to “0.1520833333333333” or “3:39”, and “03:01:02” to “3:01:02”, all of which erroneously represent times instead of alleles. The range of potential human-generated transcription

errors is large. Previous workshops devoted considerable manual effort to review, identify and correct errors, and standardize allele-name notations prior to analysis. However, the analysis, collection, exchange and storage of NGS genotyping data requires machines (computers) that are able to process allele name data, and the accompanying nucleotide sequence data, without the human ability to identify and correct errors.

1.2.4. Describing Novel Polymorphism—The description of previously unknown (novel) HLA sequence variants has been a long-standing challenge for the H&I community. Until a novel sequence is named by the World Health Organization Nomenclature Committee for Factors of the HLA System (Nomenclature Committee)[37], it is very difficult to discuss that sequence in the context of the HLA nomenclature. The common practice, associated with pre-NGS genotyping, has been to append a “novel-allele” identifier to a truncated version of a related allele name (e.g. “HLA-A*02V”, “HLA-A*02:NEW”, “HLA-A*02:01new”, etc.). The World Marrow Donor Association guidelines for the use of HLA nomenclature (WMDA guidelines) indicate that “NEW” should be reported for alleles that have not been named by the Nomenclature Committee[38]. However, the absence of a standard for describing novel HLA alleles and associated nucleotide sequences represents a considerable challenge for the collection of NGS HLA genotyping data.

2. Meeting the Challenge

The 17th IHIW adopted a centralized data-storage approach, in which all specimen-related data, reference data, genotyping data and associated meta-data were stored in a single database system. The goal of this effort was to facilitate data and analysis access for workshop participants, with these workshop products and the database itself made available to the H&I community after the workshop. The 17th IHIW focus on NGS provided an advantage for centralized data collection in that there are currently only a small number NGS platforms, which generate primary data in FASTQ[39] format, and associated genotyping software. A key 17th IHIW goal was to collect machine-generated HLA data for consumption by IHIW informatics services, with minimal human intervention. We worked with NGS software developers to define a small number of equivalent and interchangeable data reporting formats that allowed genotyping data and meta-data to be collected using a “uniform NGS data-collection” approach. This approach built on the genotype list (GL) string format[40] and the GL Service[41], the Minimum Information for Reporting Immunogenomic NGS Genotyping (MIRING) reporting guidelines and messaging standard[42], and the MIRING-compliant Histoimmunogenetics Markup Language (HML) version 1.0 messaging format[43].

2.1. Uniform NGS Data Collection

The 17th IHIW did not require all workshop projects or participating laboratories to use the same NGS platform, typing kit or protocol. NGS instruments manufactured by Illumina (e.g., MiSeq), One Lambda (e.g., S5XL), Pacific Biosciences (e.g., PacBio RSII) and Roche 454 (e.g., GS FLX) were used in 17th IHIW NGS genotyping efforts. The goal in uniform NGS data collection was that all NGS HLA genotyping data and associated meta-data (which together constitute a “typing report”) be compatible and comparable, so that all

collected data were equally interpretable, regardless of the format in which those data were exchanged. This allowed data generated by different laboratories, in different countries, using different platforms and software, to be stored in one database and made available for multiple projects.

Toward this end, the 17th IHIW accepted NGS genotyping data and meta-data in three MIRING-compliant eXtensible Markup Language (XML)[44] based typing report document formats – HML (version 1.0.1); GenDx XML, exported by GenDx NGS Engine version 2.4.0; and IHIW XML^A, a format developed specifically for the 17th IHIW (Supplements A and B). HML was generated by HistoGenetics, Omixon HLA Twin (version 1.1.4.2), Immucor MIA FORA (version 3.1) and One Lambda TypeStream Visual (version 1.1) software. IHIW XML typing reports were generated using the 17th IHIW Database (WS Database) system (described in section 2.2), by individual laboratories (using the Supplementary Materials), and by Illumina, using a “.cgp” file exported by TruSight HLA Assign version 2.1 RUO. We were unable to define a specific typing report document format for data generated on Pacific Biosciences instruments, but presumably such data could have been reported in HML or IHIW XML format. In addition, the WS Database accepted HLA genotypes in a comma-separated values (CSV) file generated by Scisco Genetics.

GenDx XML, HML and IHIW XML typing reports include subsets of the NGS genotyping data and meta-data elements described in Table 1. These data-elements are equivalent to MIRING elements 1–8[42]. HML or GenDx XML typing reports may include additional data, but because these document formats include equivalent 17th IHIW data-elements, all submitted HML and GenDx XML HLA typing reports were converted into IHIW XML typing reports (described in section 2.2.1), which were subsequently stored in the WS Database. In addition to these typing reports, the primary FASTQ data, too large to include in a report, were stored on a secure File Transfer Protocol (sFTP) server linked to the WS Database.

2.2. 17th IHIW Database

The WS Database included an Oracle SQL database (12c Standard Edition) and a web application built with APEX 5.0, running on a multi-core Linux CentOS 6 platform with 960GB of storage, expandable up to 3TB. The 17th IHIW sFTP server was an IBM high-performance computing cluster running Linux RedHat 6, with a 1Gbps Ethernet connection. The server comprised a management node, three compute nodes, two storage nodes and 15 TB of storage. The database and archived redo logs were backed up to the sFTP server with Oracle Recovery Manager (RMAN) on a daily basis, with each backup maintained for five days. The WS Database schema is illustrated in Supplementary Figure S1. WS Database tools and services were scripted in the Perl, R or Python programming languages. Both the database and the sFTP server were housed in the high-performance computing Stanford Data Center facility on the Stanford Linear Accelerator Center campus, and were managed by the Stanford Research Computing staff.

The WS Database’s structure reflected the workshop’s organization and the defined roles of workshop participants. Each of the six 17th IHIW components – NGS of HLA, NGS of KIR, Hematopoietic Cell Transplantation, Mapping of Serologic Epitopes, Informatics of

Genomic Data, and Quality Control & Quality Assurance – was led by a Component Chair (or Chairs). Projects were associated with each component, with a Principal Investigator (PI) for each project. PIs enrolled Lab Members, and could enroll in Components and Projects. Lab Members uploaded and managed data, and could enroll as Project and Component Affiliates. Further details of these participant roles can be found online^B.

The WS Database system^C stored data from typing reports and Scisco CSV tables, FASTQ files, subject and specimen data, and pedigrees (PED format[45]), and managed the accounts and data-access privileges for 17th IHIW principal investigators and lab members, project leaders, and component affiliates and chairs. When laboratory-initiated subject IDs were submitted to the WS Database, those IDs were anonymized and linked to unique 17th IHIW IDs, which were used to identify those subjects in genotyping and analysis efforts, to avoid the distribution of protected health information. The WS Database also stored project-specific data, using custom document formats, and analytic results.

2.2.1. Participant Initiated Management of Typing Reports—The submission and management of typing reports is illustrated in Figure 1. Genotyping data and meta-data could be manually entered into the WS Database, and laboratory-generated and Illumina IHIW XML typing reports were submitted directly to the WS Database. HML and GenDx XML typing reports were converted to IHIW XML reports by uploading them to the sFTP server, and using the WS Database tools to generate IHIW XML reports from them. These converted IHIW XML typing reports were stored in the WS Database, where they were available for download by participants. Regardless of their source, all IHIW XML typing reports were submitted to and stored in the WS Database. Detailed instructions on the 17th IHIW data-submission process are available online^D.

2.3. 17th IHIW Standards and Tools

To facilitate uniform NGS data collection for the 17th IHIW, we have adopted specific data-standards and conventions for the validation of typing reports, and the analysis of workshop data. The tools described in sections 2.3.1 to 2.3.4 are available on GitHub^{E,F}.

2.3.1. Typing Report Validation—Given the number of typing report formats accepted by the WS Database, we developed a number of tools and services for validating the format and content of each. Several of these tools were built into the WS Database, and ran when typing reports were uploaded or created in the system. The semantic validations and WS Database functions applied to each typing report format are listed in Table 2. Because HML and GenDx XML typing reports were converted into IHIW XML reports, the validation and database functions listed for IHIW XML format were applied to all typing reports. In addition, software developers generating HML typing reports were encouraged to use the public MIRING validator for HML service (miring-validator^E) as part of their development efforts. This validator determines if a potential HML typing report follows basic HML and MIRING rules of syntax, and if it contains MIRING dataelements. Because this validator operates as a web-service, it can be built into an HML typing report generation pipeline.

2.3.2. IPD-IMGT/HLA Database Versions—As noted in section 1.2.1, the Reference Database is updated quarterly; the number of alleles increases with each release, and the extent of sequence known for a given allele, as well as the number of fields in a given allele name, can increase between database releases. We addressed this by “freezing” all WS Database functions at Reference Database version 3.25.0 (released July 2016). While HLA allele names described in other Reference Database versions could have been submitted using HML or GenDx XML, the WS Database translated those names to their 3.25.0 counterparts upon submission (described in section 2.3.4), and all HLA-related data were analyzed using Reference Database version 3.25.0, which was also the source of all reference allele sequences. Restricting the WS Database to a single Reference Database version in this way streamlined database functions, facilitated uniform data management and analysis, and will allow the final WS Database product to be updated to later Reference Database versions in future workshops. The Reference Database resources described below are available from the Reference Database FTP site^G.

To facilitate the use of Reference Database 3.25.0 for the 17th IHIW, we defined a set of full-length (“genomic”) version 3.25.0 reference alleles (Table 3) for use in generating and aligning consensus sequences, and identifying novel polymorphism. Though some alleles in this set have names with fewer than four fields, indicating that no synonymous or non-coding polymorphism has been identified for those alleles as of Reference Database version 3.25.0, genomic sequence was available for all of them. When possible, a reference allele was identified for each allele family at a locus, but for some loci only a single full-length reference allele was identified.

2.3.3. Genotype Format and Validation—The large variety of formats used in the H&I community to record HLA genotypes and describe typing ambiguity made it difficult to collect genotyping data in a uniform manner. We addressed this by collecting all HLA genotypes in GL string format[40], using the strict-mode GL Service[41] to validate the allele content of the GL string, and applying python scripts (pyglstring^E) to validate the structure of the GL string. Data submitters were notified when GL strings failed validation (see section 2.3.5.2.1), and were requested to modify them accordingly.

These structural validation scripts we applied include an exception for the DRB3, DRB4 and DRB5 loci (the secondary DRB loci), permitting combinations of alleles at these loci to be connected by the GL string “+” operator (e.g., “HLA-DRB3*01:26N+HLA-DRB5*01:01:01”), whereas for other loci, the “+” operator connects only alleles of a single locus. When GL string format was introduced[40], the “+” operator denoted the number of copies of a given gene present in an individual. Ideally, given the structural haplotype variation known for the DRB loci[46], when the absence of a DRB3, DRB4 or DRB5 locus can be determined, the absence of that locus should be noted in a GL string. The WMDA guidelines indicate that the absence of any allele at a secondary DRB locus be reported using “NNNN” (e.g., “HLADRB1* NNNN”)[38], but this is not a widely used approach. Without a standard nomenclature for describing the confirmed absence of a secondary DRB gene, we treat these loci as alleles of a single locus. The development of a nomenclature for describing the confirmed absence of a locus (e.g. “HLA-DRB3* NNNN”, “HLA-

DRB3*00:00” or “HLA-DRB3*ABSENT”) should be considered by the Nomenclature Committee.

2.3.4. LiftOver Tool—As typing reports were accepted into the WS Database, HLA genotypes identified under Reference Database versions other than 3.25.0 were translated to their 3.25.0 counterparts via a LiftOver tool (IHIW17LiftOver.pm^F). Non-3.25.0 alleles are translated on the basis of their Reference Database accession numbers, as related in the Allelelist_history.txt file^G. In cases of alleles named after version 3.25.0 (e.g., HLA-A*01:01:01:05, identified in Reference Database version 3.27.0), the submitted allele name is translated to either the lowest-numbered 3.25.0 allele name with the greatest number of matching lower-order fields to the submitted allele (e.g., HLA-A*01:01:01:01 is chosen to replace HLA-A*01:01:01:05), or the reference allele for that locus (Table 3) when there are no matching lower-order fields, and the submitted allele is noted in the “Novelpolymorphism” field for that genotype (e.g. as “IPD-IMGT/HLA-3270-HLA-A*01:01:01:05”). We note that it remains unclear if a typing system that e.g., identified HLA-A*01:01:01:05 under Reference Database version 3.27.0 would return a result of HLA-A*01:01:01:01 for the given specimen under version 3.25.0. However, we felt that storage of the submitted allele name in the Novelpolymorphism field would be sufficient to allow these instances to be investigated. In cases where allele name changes that occurred in Reference Database versions prior to 3.25.0 resulted in accession number changes (e.g., HLA-DRB1*08:01:03, with accession number HLA02257, was changed to HLA-DRB1*08:01:01, with accession number HLA00723, as part of Reference Database version 3.24.0, as detailed in Table 4), the version 3.25.0 allele name was used.

When ambiguous HLA genotypes were submitted, the LiftOver tool evaluated ambiguous alleles (delimited with the GL string slash [/] operator) and ambiguous genotypes (delimited with the GL string pipe [|] operator), and identified alleles and genotypes that could be translated to their 3.25.0 counterparts (Figure 2). These alleles were translated, and the GL string consolidated to eliminate duplications. If an ambiguous HLA genotype consisted entirely of alleles named after Reference Database version 3.25.0, the LiftOver tool translated those alleles to the corresponding lowest-numbered 3.25.0 alleles with the greatest number of matching lower-order fields, as described above, and consolidated the GL string. In all cases, the submitted non-3.25.0 GL strings were stored in the “Original_GL” field for that genotype. These allelic and GL string LiftOver functions were accomplished using a modified version of the Allelelist_history.txt file that includes data from the hla_nom.txt^G files and Table 3 (IHIW17_AllelelistGgroups_history.txt^F). This LiftOver process occurred when HML and GenDx XML typing reports were converted to IHIW XML reports. All collected IHIW XML typing reports corresponded to version 3.25.0. However, the LiftOver tool can be modified to allow all collected typing reports to be updated to a future Reference Database version.

2.3.5. 17th IHIW Database Tools and Functions—Reference Database version 3.25.0 included 12.9 million bases of sequence for 14,957 HLA alleles at 19 HLA loci. Of this, more than 40,000 exons comprised 9 million bases of sequence, making this a rich, but very complex, data resource. We developed user-facing front end tools to assist 17th IHIW

participants in working with these data, and data-facing back end tools to facilitate the integration of the large quantities of new sequence that were be generated via NGS.

2.3.5.1. Front End Tools

2.3.5.1.1. *hlaPoly*: The absence of a standard method for describing novel nucleotide polymorphism in consensus sequences posed challenges for our uniform data collection approach. Typing reports generated for the same specimen using different genotyping software could include identical consensus sequences and genotypes, but when a consensus sequence includes novel polymorphism, the Reference Database version, reference allele sequence, and sequence coordinate system used to describe that polymorphism could vary between software applications, and typing reports generated by different software may have identified different novel polymorphism for identical consensus sequences. For example, the nucleotide sequence of the HLA-A*01:01:01:05 allele differs from the HLA-A*01:01:01:01 allele in Reference Database 3.25.0 at three intron 2 nucleotide positions, and differs from the HLA-A*01:01:01:03 allele in Reference Database 3.25.0 at those same three positions as well as at an intron 1 position; the reference allele used to align the HLA-A*01:01:01:05 consensus sequence informed the description of novel polymorphism.

To standardize novel polymorphism description for the 17th IHIW, we developed the *hlaPoly* R package^F, which identifies novel polymorphism for a given consensus sequence, when provided with the closest matching allele name (which was usually included in the genotype) and the Reference Database version (version 3.25.0). The *hlaPoly* tool was deployed online as a Shiny application^H. As illustrated in Supplementary Figure S2, *hlaPoly* uses the DECIPHER R package[47] to generate a multiple sequence alignment for the full-length HLA reference allele sequence (Table 3), the sequence of the pertinent allele in the genotype (closest allele) and the consensus sequence, and then retrieves the mismatches and indels between the consensus sequence and the called allele as novel polymorphism. If no sequence is known for the called allele in an aligned region, the mismatches and indels between the consensus sequence and the full-length HLA reference allele are retrieved. For each novel polymorphism, the feature number and start/end position relative to that feature are also calculated. The WS Database stored these novel polymorphism data in both a tabular form (see the bottom of Figure S1) and a string format (described in Supplement A).

2.3.5.1.2. *Quick Calculation of Feature Position*: To assist in manual entry of genotyping data and meta-data into the WS Database, we developed a tool for the calculation of gene-feature information. Given an allele name and the nucleotide position relative to the start of known nucleotide sequence for that allele, this tool returned the feature name (e.g. Exon 2), feature ID and the relative nucleotide position in that feature. This tool was available in the WS Database under “Lab Member”/”Tools”/”IMGT/HLA Feature List”.

2.3.5.1.3. *Concatenate HML files*: Each HML file uploaded to the sFTP server was treated as single typing report, and as suggested in Figure 1, some HML typing reports were generated for individual samples. Rather than requiring that hundreds or thousands of individual-sample HML files be converted to IHIW XML files, each of which would need to be manually loaded into the WS Database, we provided a tool (*concahtml.pl*^F) that

concatenates multiple HML files into a single HML file, which can be converted into a single IHIW XML file for loading. This “Concat HML files” tool was available in the WS Database under “Lab Member”/“Tools”.

2.3.5.1.4 “Convert HML to IHIW XML” and “Convert GenDX XML to IHIW XML”: As noted in Figure 1, the sFTP server automatically generated an IHIW XML typing report when an HML report was loaded into the /hml directory. The server also generated an IHIW XML report when a GenDX XML report was loaded into the /gendx directory. The “Convert HML to IHIW XML” and “Convert GenDX XML to IHIW XML” tools could be used to force these automatic functions to run immediately, or to manually convert HML or GenDX XML typing reports that had been loaded into other directories. Both tools were available in the WS Database under “Lab/Member”/“Tools”.

2.3.5.2. Back End Tools

2.3.5.2.1. Watcher Daemons: To monitor activity on the sFTP server, we developed daemons that detected new HML and GenDX XML files as they were uploaded to the sFTP server, automatically converted them to IHIW XML files, and validated them during the conversion. Any validation errors were logged and made available under “Lab Member”/“Tools”/“Job Log” in the WS Database. A second set of daemons performed daily checks for new typing reports in the WS Database. These daemons ran hlaPoly for newly added or edited consensus sequences, and stored the novel polymorphism results in the WS Database.

2.3.5.2.2. Consensus Linking: Genotypes and consensus sequences are recorded separately in HML typing reports. Each consensus sequence is associated with the reference allele used to align it, which is usually a full-length allele, but is not directly linked to specific alleles in the associated genotype. For cases when these reference alleles are not included in the genotype, we developed a consensus linking tool that identified the allele in the genotype that most closely matched the reference allele, using the same approach applied for the LiftOver process (described in section 2.3.4). For example, if HLA-A*11:01:01:01 and *31:01:02:01 are the respective reference alleles for consensus sequences A and B, which are associated with the HLA-A*11:01:28+HLA-A*31:01:07, the consensus linking tool would associate HLA-A*11:01:28 with consensus sequence A and *31:01:07 with consensus sequence B.

2.4. Support for 17th IHIW Projects

In addition to its collection, validation and storage functions, the WS Database supported 17th IHIW projects by integrating tools for HLA data analysis and exchange. An updated version of PyPop[48]^I, supporting colon-delimited allele names, with increased multi-locus analysis capacity, was accessible through the WS Database system. Similarly, integration of Gene Feature Enumeration[49] (GFE) functions (e.g., the feature-service^{E,J}, GFE service^{E,K} and Allele-Calling Tool^{E,L}) allows full-gene HLA sequences to be exchanged and analyzed in the absence of an HLA allele name.

2.5. Validation of Tools and Functions

The data collections tools and functions described in section 2.3 were validated iteratively, as part of the process of working with NGS vendors to develop each typing report format. As new submitters provided data (e.g. a new vendor generating HML, or a new laboratory generating IHIW XML) the initial data imported to the WS Database were compared manually via comparison with the submitted typing reports; in instances of discrepancies, either feedback was provided to the submitter, or the tools and functions were updated to accommodate the “new” typing report format.

2.5.1 Genotyping Expectations—In 2014, a panel of 50 IHWG reference cell lines that had been well-characterized using pre-NGS HLA-A, -B, -C, -DPA1, -DPB1, -DQA1, -DQB1, -DRB1, -DRB3, -DRB4 and -DRB5 typing methods was assembled for a 17th IHIW NGS genotyping pilot project. HLA genotyping for this pilot project cell panel was independently performed by 15 laboratories using multiple NGS methods. This set of 15 NGS genotyping results was used to establish concordance expectations for the initial quality control (QC) and ongoing proficiency testing (PT) required for all 17th IHIW participants performing NGS genotyping. Random subsets of 24 pilot project cell lines were sent to each genotyping laboratory for QC/PT, and we used these QC/PT results to validate the genotype-data manipulations and transformations performed by the tools, functions and services described in section 2.3.

2.5.2 Manual Validation of Performance—Parsers and converters that did not depend on algorithmic transformations were validated by manual inspection of the results. For example, during hlaPoly development, output tables were validated manually, by comparison with novelPolymorphism results included in typing reports. The LiftOver process was validated using the typing reports generated by each laboratory as part of the QC/PT process.

3. Conclusions

We addressed several of the long-standing challenges to uniform NGS HLA data-collection and -storage for the 17th IHIW by developing new tools and formats, and adopting existing standards and services. NGS vendors worked with us to develop equivalent NGS HLA typing report formats that ensured data-portability across 17th IHIW projects. We ensured data-quality by validating all typing reports before they were loaded to the WS Database. All HLA genotyping data were recorded using the same Reference Database version, and novel HLA polymorphism was described using the same reference alleles. This approach facilitated the basic and clinical research aims of 17th IHIW HLA Projects, and can be applied by the larger H&I community. The 17th IHIW was held in September of 2017. We hope to work with the organizers of the 18th IHIW so that the WS Database and its associated tools will serve as a persistent central H&I community resource that will ensure research and data continuity with future IHIW efforts.

Supplementary Materials

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This work was supported by National Institutes of Health (NIH) National Institute of Allergy and Infectious Disease (NIAID) grant R01AI128775 (BM, MM, SM), NIH National Institute of General Medical Sciences (NIGMS) grant R01GM109030 (BM, MM, SM), Office of Naval Research (ONR) grant N00014-08-1-1207 (BM and MM), and an overseas project funded by the Taiwanese Ministry of Science and Technology (MST) (CC). The content is solely the responsibility of the authors and does not necessarily reflect the official views of the MST, NIAID, NIGMS, NIH, ONR, Taiwanese government or United States government. We thank the Stanford Blood Center for the support and promotion of the 17th IHIWS endeavor, Ken Yamaguchi for helpful discussions and manuscript review, Tamara Vayntrub for her tremendous administrative support of 17th IHIW efforts, and the histocompatibility and immunogenetics community and the International HLA and Immunogenetics Workshop Council for their continued dedication to and support of the International Workshops. We also thank President Barack H. Obama for his support and appreciation of American science and basic research; several of the authors owe their continuing scientific careers to the passage of the American Recovery and Reinvestment Act (ARRA), and while ARRA funds did not support this work, it would not have been possible without ARRA.

Abbreviations

CSV	Comma-Separated Values
GFE	Gene Feature Enumeration
GL	Genotype List
HLA	Human Leukocyte Antigen
HML	Histoimmunogenetics Markup Language
H&I	Histocompatibility and Immunogenetics
IHIW	International HLA and Immunogenetics Workshop
IMGT	ImMunoGeneTics
IPD	ImmunoPolymorphism Database
IUPAC	International Union of Pure and Applied Chemistry
KIR	Killer-cell Immunoglobulin-like Receptor
MIRING	Minimum Information for Reporting Immunogenomic NGS Genotyping
NGS	Next Generation Sequencing
PI	Principal Investigator
RMAN	Recovery Manager
RSCA	Reference Strand Conformation Analysis
rSSO	Reverse Sequence-Specific Oligo
SBT	Sequence-Based Typing
sFTP	secure File Transfer Protocol
SS	Sequence-Specific

SSO	Sequence-Specific Oligo
SSP	Sequence-Specific Priming
WMDA	World Marrow Donor Association
WS	Workshop
XML	eXtensible Markup Language

Literature Cited

1. Scandinavian Journal of Haematology; Histocompatibility Testing 1965: Report of a Conference and Workshop Sponsored by the Boerhaave Courses for Postgraduate Medical Education, University of Leiden; Aug 15–21, 1965; Copenhagen: Munksgaard; 1965.
2. Histocompatibility Testing; Report of a Conference and Workshop Sponsored by the Division of Medical Sciences, National Academy of Sciences, National Research Council; 7–12 June, 1964; Washington: National Academy of Sciences; 1965.
3. Curtoni, ES., Mattiuz, PL., Tosi, RM. Histocompatibility Testing 1967. Copenhagen: Munksgaard; 1967.
4. Terasaki, PI. Histocompatibility Testing 1970. Copenhagen: Munksgaard; 1970.
5. Dausset, J., Colombani, J. Histocompatibility Testing 1972. Copenhagen: Munksgaard; 1973.
6. Kissmeyer-Nielsen, F. Histocompatibility Testing 1975. Copenhagen: Munksgaard; 1975.
7. Bodmer, WF., Batchelor, JR., Bodmer, JG., Festenstein, H., Morris, PJ. Histocompatibility Testing 1977. Copenhagen: Munksgaard; 1978.
8. Terasaki, PI. Histocompatibility Testing 1980. Los Angeles: UCLA Tissue Typing Laboratory; 1980.
9. Albert, ED., Baur, MP., Mayer, WR. Introductory Remarks. In: Albert, ED., Baur, MP., Mayer, WR., editors. Histocompatibility Testing 1984. Heidelberg: Springer-Verlag; 1984.
10. Dupont, B. Overview of Experimental Design for the Tenth International Histocompatibility Workshop. In: Dupont, B., editor. Immunobiology of HLA. Volume I. Histocompatibility Testing. Vol. 1. Springer Verlag; 1987. p. 3
11. Tsuji, K. Overview of the Eleventh International Histocompatibility Workshop and Conference. In: Kimiyoshi Tsujiki, MA., Sasazuki, Takehiko, editors. HLA 1991: Proceedings of the Eleventh International Histocompatibility Workshop and Conference. Vol. 1. Oxford Science Publications; 1991. p. 3
12. Charron, D., Fauchet, R. The 12th International Histocompatibility Workshop. In: Charron, D., editor. HLA Volume 1. Genetic Diversity of HLA: Functional and Medical Implication. Vol. 1. EDK; 1997. p. XXV
13. Hansen, JA. Foreword: Immunobiology of the Human MHC. Proceedings of the 13th International Histocompatibility Workshop and Conference. In: Hansen, JA., editor. Immunobiology of the Human MHC: Proceedings of the 13th International Histocompatibility Workshop and Conference. Volume 1. Vol. 1. IHWG Press; 1997. p. xxv
14. McCluskey J, Tait CB, Christiansen FT, Holdsworth R. 14th International HLA and Immunogenetics Workshop Reports: Introduction. Tissue Antigens. 2007; 69:1. [PubMed: 17212702]
15. Tissue Antigens; Abstracts for the 15th International Histocompatibility and Immunogenetics Workshop and Conference; Rio de Janeiro, Brazil. September 13–20, 2008; 2008. p. 231
16. Middleton D, Marsh SGE. 16th International HLA and Immunogenetics Workshop (IHIW) Introduction. Int J Immunogenet. 2013; 40:1. [PubMed: 23280276]
17. Baur, MP., Albert, ED., Mayr, WR. The Central Data Analysis of the Ninth Workshop. In: Albert, ED., Baur, MP., Mayr, WR., editors. Histocompatibility Testing 1984. Heidelberg: Springer-Verlag; 1984. p. 37

18. Lalouel, J-M., Ferguson, M., Wheeler, R., King, N. Tenth International Histocompatibility Workshop: Overview of Data Processing and Analysis. In: Dupont, B., editor. Immunobiology of HLA. Volume I. Histocompatibility Testing 1987. Vol. 1. Springer-Verlag; 1987. p. 83
19. Inoko, H., Sato, K., Takata, H., Imanishi, T., Ina, Y., Saitou, N., et al. Data Flow from the Collection of Raw Data to the Construction of the MHC Database. In: Kimiyoshi Tsujiki, MA., Sasazuki, Takehiko, editors. HLA 1991. Proceedings of the Eleventh International Histocompatibility Workshop and Conference. Vol. 1. Oxford Science Publications; 1991. p. 65
20. Cambon-Thomsen, A., Albert, E., Bodmer, JG., Piazza, A., Thouzellier, Y., Clayton, JF. Database, Communication, Analysis (DCA) of the 12th International Histocompatibility Workshop: General Overview. In: Charron, D., editor. HLA Volume 1. Genetic Diversity of HLA: Functional and Medical Implications. Vol. 1. EDK; 1997. p. 469
21. Schoch, G., Kesten, M., McKallor, C., Mickelson, E., Hansen, JA. 13th IHWS Shared Resources Joint Report. In: Hansen, JA., editor. Immunobiology of the Human MHC: Proceedings of the 13th International Histocompatibility Workshop and Conference. Volume I. Vol. 1. IHWG Press; 2007. p. 482
22. Nunes JM. Tools for analysing ambiguous HLA data. *Tissue Antigens*. 2007; 69:203.
23. Mack SJ, Sanchez-Mazas A, Single RM, Meyer D, Hill J, Dron HA, et al. Population samples and genotyping technology. *Tissue Antigens*. 2007; 69:188. [PubMed: 17445198]
24. Single RM, Meyer D, Mack SJ, Lancaster A, Erlich HA, Thomson G. 14th International HLA and Immunogenetics Workshop: Report of progress in methodology, data collection, and analyses. *Tissue Antigens*. 2007; 69:185. [PubMed: 17445197]
25. Steenkiste A, Valdes AM, Feolo M, Hoffman D, Concannon P, Noble J, et al. 14th International HLA and Immunogenetics Workshop: Report on the HLA component of type 1 diabetes. *Tissue Antigens*. 2007; 69:214. [PubMed: 17445204]
26. Leffell MS, Cao K, Coppage M, Hansen JA, Hart JM, Pereira N, et al. Incidence of humoral sensitization in HLA partially mismatched hematopoietic stem cell transplantation. *Tissue Antigens*. 2009; 74:494. [PubMed: 19804563]
27. Middleton D, Gonzalez F, Fernandez-Vina M, Tiercy JM, Marsh SGE, Aubrey M, et al. A bioinformatics approach to ascertaining the rarity of HLA alleles. *Tissue Antigens*. 2009; 74:480. [PubMed: 19793314]
28. Hollenbach JA, Meenagh A, Sleator C, Alaez C, Bengoche M, Canossi A, et al. Report from the killer immunoglobulin-like receptor (KIR) anthropology component of the 15th International Histocompatibility Workshop: worldwide variation in the KIR loci and further evidence for the co-evolution of KIR and HLA. *Tissue Antigens*. 2010; 76:9. [PubMed: 20331834]
29. Nunes JM, Riccio ME, Buhler S, Di D, Currat M, Ries F, et al. Analysis of the HLA population data (AHPD) submitted to the 15th International Histocompatibility/Immunogenetics Workshop by using the Gene[rate] computer tools accommodating ambiguous data (AHPD project report). *Tissue Antigens*. 2010; 76:18. [PubMed: 20331842]
30. Naumova, E., Ivanova, M., Pawelec, G., Constantinescu, I., Bogunia-Kubik, K., Lange, A., et al. *Tissue Antigens*; 'Immunogenetics of Aging': report on the activities of the 15th International HLA and Immunogenetics Working Group and 15th International HLA and Immunogenetics Workshop; 2011. p. 187
31. Riccio ME, Buhler S, Nunes JM, Vangenot C, Cuénod M, Currat M, et al. 16th IHIW: Analysis of HLA Population Data, with updated results for 1996 to 2012 workshop data (AHPD project report). *Int J Immunogenet*. 2013; 40:21. [PubMed: 23280239]
32. Gonzalez-Galarza FF, Mack SJ, Hollenbach J, Fernandez-Vina M, Setterholm M, Kempenich J, et al. 16th IHIW: Extending the number of resources and bioinformatics analysis for the investigation of HLA rare alleles. *Int J Immunogenet*. 2013; 40:60. [PubMed: 23198982]
33. Hollenbach JA, Mack SJ, Gourraud PA, Single RM, Maiers M, Middleton D, et al. A community standard for immunogenomic data reporting and analysis: proposal for a STrengthening the REporting of Immunogenomic Studies statement. *Tissue Antigens*. 2011; 78:333. [PubMed: 21988720]
34. Robinson J, Soormally AR, Hayhurst JD, Marsh SG. The IPD-IMGT/HLA Database - New developments in reporting HLA variation. *Hum Immunol*. 2016; 77:233. [PubMed: 26826444]

35. Marsh SG, Albert ED, Bodmer WF, Bontrop RE, Dupont B, Erlich HA, et al. Nomenclature for factors of the HLA system, 2002. *Tissue Antigens*. 2002; 60:407. [PubMed: 12492818]
36. Marsh SG, Albert ED, Bodmer WF, Bontrop RE, Dupont B, Erlich HA, et al. Nomenclature for factors of the HLA system, 2010. *Tissue Antigens*. 2010; 75:291. [PubMed: 20356336]
37. Committee WN Nomenclature for factors of the HL-a system. *Bull World Health Organ*. 1968; 39:483. [PubMed: 5303912]
38. Bochtler W, Maiers M, Oudshoorn M, Marsh SG, Raffoux C, Mueller C, et al. World Marrow Donor Association guidelines for use of HLA nomenclature and its validation in the data exchange among hematopoietic stem cell donor registries and cord blood banks. *Bone Marrow Transplant*. 2007; 39:737. [PubMed: 17438587]
39. Cock PJ, Fields CJ, Goto N, Heuer ML, Rice PM. The Sanger FASTQ file format for sequences with quality scores, and the Solexa/Illumina FASTQ variants. *Nucleic Acids Res*. 2010; 38:1767. [PubMed: 20015970]
40. Milius RP, Mack SJ, Hollenbach JA, Pollack J, Heuer ML, Gragert L, et al. Genotype List String: a grammar for describing HLA and KIR genotyping results in a text string. *Tissue Antigens*. 2013; 82:106. [PubMed: 23849068]
41. Milius RP, Heuer M, George M, Pollack J, Hollenbach JA, Mack SJ, et al. The GL service: Web service to exchange GL string encoded HLA & KIR genotypes with complete and accurate allele and genotype ambiguity. *Hum Immunol*. 2016; 77:249. [PubMed: 26621609]
42. Mack SJ, Milius RP, Gifford BD, Sauter J, Hofmann J, Osoegawa K, et al. Minimum information for reporting next generation sequence genotyping (MIRING): Guidelines for reporting HLA and KIR genotyping via next generation sequencing. *Hum Immunol*. 2015; 76:954. [PubMed: 26407912]
43. Milius RP, Heuer M, Valiga D, Doroschak KJ, Kennedy CJ, Bolon YT, et al. Histoimmunogenetics Markup Language 1. 0: Reporting next generation sequencing-based HLA and KIR genotyping. *Hum Immunol*. 2015; 76:963. [PubMed: 26319908]
44. Bray T, Paoli J, Sperberg-McQueen CM, Maler E, Yergeau F. Extensible markup language (XML). World Wide Web Consortium Recommendation REC-xml-19980210. 1998; 16:16. <http://www.w3.org/TR/1998/REC-xml-19980210>.
45. Purcell S, Neale B, Todd-Brown K, Thomas L, Ferreira MA, Bender D, et al. PLINK: a tool set for whole-genome association and population-based linkage analyses. *Am J Hum Genet*. 2007; 81:559. [PubMed: 17701901]
46. Andersson G. Evolution of the human HLA-DR region. *Front Biosci*. 1998; 27:d739.
47. Wright ES. Using DECIPHER v2. 0 to analyze big biological sequence data in R. *The R Journal*. 2016; 8:352.
48. Lancaster AK, Single RM, Nelson MP, Solberg O, Thomson G. PyPop update - a software pipeline for large-scale multi-locus population genomics. *Tissue Antigens*. 2007; 69:192. [PubMed: 17445199]
49. Mack SJ. A gene feature enumeration approach for describing HLA allele polymorphism. *Hum Immunol*. 2015; 76:975. [PubMed: 26416087]
50. Cornish-Bowden A. Nomenclature for incompletely specified bases in nucleic acid sequences: recommendations 1984. *Nucleic Acids Res*. 1985; 13:3021. [PubMed: 2582368]

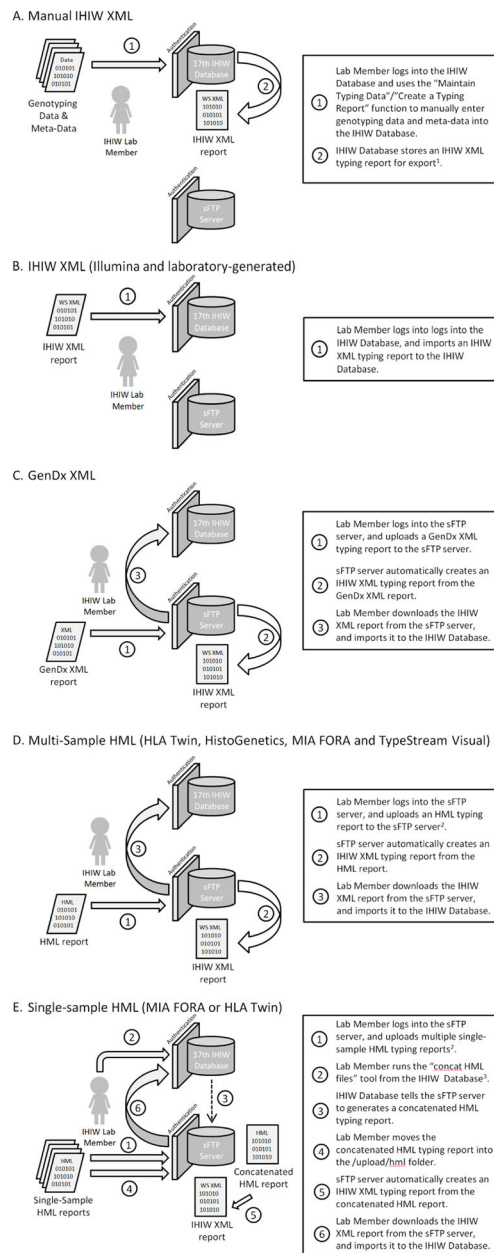


Figure 1. Typing Report Submission Procedures

Cartoon descriptions describing the key steps of the five 17th IHIW typing report submission processes are shown, with the steps defined in each inset box.

HML: Histoimmunogenetics Markup Language

IHIW: International HLA and Immunogenetics Workshop

sFTP: secure File Transfer Protocol

XML: eXtensible Markup Language

1: Though this final step is only shown in panel A, an IHIW XML typing report remains available for download from the WS Database after an IHIW XML report of any source is loaded into the WS Database.

2: WS Database watcher daemons monitor the sFTP server's */upload/hml* directory for the arrival of HML typing reports, and trigger the automatic conversion HML reports into IHIW XML reports. Multi-sample (project-level) HML reports should be loaded directly into the */upload/hml* directory. Single-sample HML reports should be loaded into a user-created subdirectory of the */upload* directory.

3: For the "concat HML files" tool to run, the Lab Member must supply the user-created subdirectory of the */upload* directory in which the single-sample HML reports have been loaded.

- A. Manual entry of genotyping data and meta-data to the WS Database
- B. Illumina-generated or laboratory-generated IHIW XML
- C. GenDx XML
- D. Multi-sample (project-level) HML generated by HistoGenetics, HLA Twin, MIA FORA and TypeStream Visual
- E. Single-sample HML generated by MIA FORA or HLA Twin

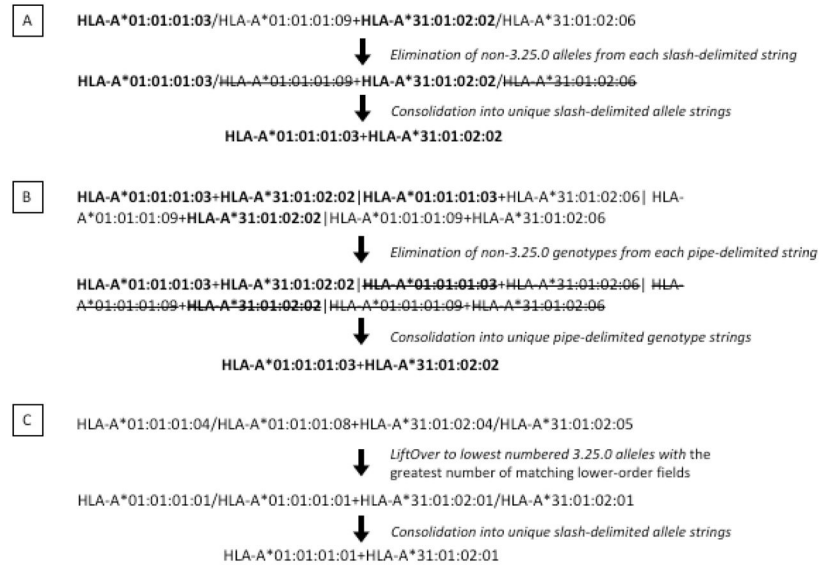


Figure 2. Example of LiftOver of Ambiguous HLA-A Genotypes

Examples of the LiftOver process for three ambiguous HLA-A genotypes consistent with Reference Database version 3.28.0 are shown. Allele names that are included in Reference Database 3.25.0 are shown in boldface.

- LiftOver process for slash-delimited ambiguous allele strings that include 3.25.0 alleles.
- LiftOver process for pipe-delimited ambiguous genotype strings that include 3.25.0 alleles.
- LiftOver process for slash-delimited allele strings that do not include 3.25.0 alleles.

Table 1

Data-Elements in 17th IHIW Typing Reports

Data-Elements	Data Type	Description	Typing Report Format ^a
17th IHIW Lab-code	Identifier	A 6-character code provided by the 17th IHIW to identify each participating laboratory.	ABCDE
Report ID	Identifier	A code provided by the originating lab to identify each report.	ABCDE
Specimen ID	Identifier	A 17th IHIW code that uniquely identifies the specimen that was genotyped.	ABCDE
Instrument	Meta-data	Parameters that document the name, manufacturer, model, and on-board software of each instrument used to generate the typing.	AB
Reagent Protocol	Meta-data	Parameters that document the name, manufacturer, and reference source for any reagents or kits used to generate the typing, along with protocol deviations.	AB
Software	Meta-data	Parameters that document the name, manufacturer and version of each program used to generate the typing, along with the use to which that program was applied, and any non-default parameters applied.	ABC
Reference Database Version	Meta-data	Documentation of the IPD-IMGT/HLA Database release version(s) used for the sequence alignment and base-calling that generated the consensus sequence and genotype.	ABCDE
Reference Sequence	Meta-data	The identifiers for the reference sequences used for the sequence alignment and base calling that generated the consensus sequence and genotype	CDE/
Locus	Genotyping data	The locus associated with each genotype and consensus sequence.	ABCDE
Genotype	Genotyping data	A genotype written in GL-String format[40] for each locus typed.	ABDE2
Consensus Sequence	Genotyping data	A nucleotide sequence representing a contiguous phased region of DNA.	ABCDE
Sequence Coordinate	Meta-data	The start and end positions of the consensus sequence(s) with respect to the reference sequence.	ABCDE
Phasing	Meta-data	Parameters that describe the phase relationships between the consensus sequences at each locus.	ABCDE
Sequence Feature	Meta-data	The gene feature or features (exons, introns or untranslated regions) represented by the consensus sequence	AB, ³
Sequence Quality	Meta-data	The mean depth of reads used to generate a given consensus sequence	AC
Typing Annotation	Meta-data	A structured notation for identifying instances when allele names included in the genotype are the closest matches to the consensus sequence, but do not correspond exactly to the reported consensus sequence.	A ⁴
Novel Polymorphism	Genotyping data	A description of any novel polymorphism detected.	ABCDE
FASTQ Location	Meta-data	The name and location (in the WS Database, or online) of the primary ("raw") FASTQ data for each genotype	ACDE

IHIW: International HLA and Immunogenetics Workshop

GL: Genotype List

IPD-IMGT: ImmunoPolymorphism Database-ImmunoGeneTics

^aFor each data-element, the typing report format in which it is found it is listed. As referenced in Figure 1. A: Manual IHIW XML; B: Illumina and laboratory-generated IHIW XML; C: GenDx XML; D: HLA Twin, HistoGenetics, MIA FORA and TypeStream Visual HML; E: HLA Twin and MIA FORA HML.

Author Manuscript

Author Manuscript

Author Manuscript

Author Manuscript

¹The A and B formats use the reference sequences in Table 3.

²The WS Database conversion daemon generates GL Strings for format C.

³The C, D, and E formats use the “Genomic - Unknown Location” sequence feature.

⁴The hlaPoly tool identifies this information for all typing report formats.

Table 2

Validation and Database Functions Applied to each typing report format

HML	Validation	<ol style="list-style-type: none"> 1 GL-String content validation with strict-mode GL service of the provided Reference Database version 2 GL-String "sanity check" syntax validation
	Functions	<ol style="list-style-type: none"> 1 GL-String LiftOver to Reference Database version 3.25.0 2 GL-String concatenation 3 Retrieve HLA typing from GL-String using the provided reference allele
GenDx	Functions	<ol style="list-style-type: none"> 1 Generate GL-String from GenDx "genotype list" elements 2 Generate phasing groups
IHIW XML/Manual Entry	Validation	<ol style="list-style-type: none"> 1 Identifier (e.g. 17th IHIW labcode, sample ID) validation 2 GL-String validation with strict-mode 3.25.0 GL service 3 The uniqueness of the quartet annotation (sample ID, HLA typing, phasing group, start position) of consensus sequences 4 IUPAC nucleotide code[50] validation for consensus sequence
	Functions	<ol style="list-style-type: none"> 1 hlaPoly application to identify novel polymorphisms

HML: Histoimmunogenetics Markup Language

IHIW: International HLA and Immunogenetics Workshop

IUPAC: International Union of Pure and Applied Chemistry

XML: eXtensible Markup Language

Because HML and GenDx XML typing reports are converted to IHIW XML reports upon submission, the validation and functions listed for IHIW WS format are applied to all typing reports.

Validation results and details are provided in the WS Database system under "Lab Member"/"Tools"/"Job Log".

Table 3

Full-length HLA Reference Alleles in IPD-IMGT/HLA Database Version 3.25.0

Locus	Accession Number	Allele Name	Description ^a
HLA-A	HLA00001	HLA-A*01:01:01:01	HLA-A Reference A*01 Reference A*36 Reference
	HLA00005	HLA-A*02:01:01:01	A*02 Reference
	HLA00037	HLA-A*03:01:01:01	A*03 Reference
	HLA00043	HLA-A*11:01:01:01	A*11 Reference
	HLA00048	HLA-A*23:01:01	A*23 Reference
	HLA00050	HLA-A*24:02:01:01	A*24 Reference
	HLA00071	HLA-A*25:01:01	A*25 Reference A*26 Reference
	HLA00085	HLA-A*29:01:01:01	A*29 Reference
	HLA00089	HLA-A*30:01:01	A*30 Reference
	HLA00097	HLA-A*31:01:02:01	A*31 Reference
	HLA00101	HLA-A*32:01:01	A*32 Reference
	HLA00104	HLA-A*33:01:01	A*33 Reference
	HLA00108	HLA-A*34:01:01	A*34 Reference
	HLA00112	HLA-A*66:01:01	A*43 Reference A*66 Reference
	HLA05918	HLA-A*68:01:01:02	A*68 Reference A*69 Reference
	HLA05527	HLA-A*74:02:01:02	A*74 Reference
HLA00130	HLA-A*80:01:01:01	A*80 Reference	
HLA-B	HLA00132	HLA-B*07:02:01	HLA-B Reference B*07 Reference B*82 Reference B*83 Reference
	HLA00146	HLA-B*08:01:01:01	B*08 Reference
	HLA00152	HLA-B*13:01:01	B*13 Reference
	HLA00157	HLA-B*14:01:01	B*14 Reference
	HLA00162	HLA-B*15:01:01:01	B*15 Reference
	HLA00213	HLA-B*18:01:01:01	B*18 Reference
	HLA00221	HLA-B*27:02:01	B*27 Reference
	HLA00237	HLA-B*35:01:01:01	B*35 Reference
	HLA00265	HLA-B*37:01:01	B*37 Reference
	HLA00267	HLA-B*38:01:01	B*38 Reference B*39 Reference
	HLA00292	HLA-B*40:01:02	B*40 Reference
	HLA13397	HLA-B*40:305	B*41 Reference
	HLA00315	HLA-B*42:01:01	B*42 Reference
	HLA00318	HLA-B*44:02:01:01	B*44 Reference
HLA00329	HLA-B*45:01:01	B*45 Reference	

Locus	Accession Number	Allele Name	Description ^a
	HLA00331	HLA-B*46:01:01	B*46 Reference
	HLA14088	HLA-B*47:01:01:03	B*47 Reference
	HLA00335	HLA-B*48:01:01	B*48 Reference B*81 Reference
	HLA00340	HLA-B*49:01:01	B*49 Reference B*50 Reference
	HLA00344	HLA-B*51:01:01:01	B*51 Reference B*52 Reference B*78 Reference
	HLA00364	HLA-B*53:01:01	B*53 Reference B*58 Reference
	HLA00367	HLA-B*54:01:01	B*54 Reference B*55 Reference B*56 Reference B*59 Reference
	HLA00381	HLA-B*57:01:01	B*57 Reference
	HLA00390	HLA-B*67:01:01	B*67 Reference
	HLA00392	HLA-B*73:01	B*73 Reference
HLA-C	HLA00401	HLA-C*01:02:01	HLA-C Reference C*01 Reference
	HLA00405	HLA-C*02:02:02:01	C*02 Reference
	HLA01543	HLA-C*03:02:02:01	C*03 Reference
	HLA00420	HLA-C*04:01:01:01	C*04 Reference
	HLA00427	HLA-C*05:01:01:01	C*05 Reference
	HLA00430	HLA-C*06:02:01:01	C*06 Reference
	HLA00433	HLA-C*07:01:01:01	C*07 Reference
	HLA00445	HLA-C*08:01:01	C*08 Reference
	HLA00454	HLA-C*12:02:02	C*12 Reference
	HLA00462	HLA-C*14:02:01:01	C*14 Reference
	HLA00467	HLA-C*15:02:01:01	C*15 Reference
	HLA00475	HLA-C*16:01:01:01	C*16 Reference
	HLA00481	HLA-C*17:01:01:01	C*17 Reference
HLA00483	HLA-C*18:01	C*18 Reference	
HLA-DPA1	HLA06604	HLA-DPA1*01:03:01:02	HLA-DPA1 Reference DPA1*01 Reference DPA1*03 Reference DPA1*04 Reference
	HLA00505	HLA-DPA1*02:01:02	DPA1*02 Reference
HLA-DPB1	HLA00517	HLA-DPB1*02:01:02	HLA-DPB1 Reference
HLA-DQA1	HLA00601	HLA-DQA1*01:01:01:01	HLA-DQA1 Reference
HLA-DQB1	HLA00622	HLA-DQB1*02:01:01	HLA-DQB1 Reference
HLA-DRB1	HLA00664	HLA-DRB1*01:01:01	HLA-DRB1 Reference DRB1*01 Reference DRB1*10 Reference
	HLA00671	HLA-DRB1*03:01:01:01	DRB1*03 Reference
	HLA00685	HLA-DRB1*04:01:01:01	DRB1*04 Reference

Locus	Accession Number	Allele Name	Description ^a
	HLA00719	HLA-DRB1*07:01:01:01	DRB1*07 Reference
	HLA00727	HLA-DRB1*08:03:02	DRB1*08 Reference
	HLA09928	HLA-DRB1*09:21	DRB1*09 Reference
	HLA00751	HLA-DRB1*11:01:01:01	DRB1*11 Reference
	HLA14829	HLA-DRB1*12:01:01:02	DRB1*12 Reference
	HLA00797	HLA-DRB1*13:01:01:01	DRB1*13 Reference
	HLA00837	HLA-DRB1*14:05:01	DRB1*14 Reference
HLA-DRB1	HLA03453	HLA-DRB1*15:01:01:02	DRB1*15 Reference DRB1*16 Reference
HLA-DRB3	HLA00887	HLA-DRB3*01:01:02:01	HLA-DRB3 Reference
HLA-DRB4	HLA00905	HLA-DRB4*01:01:01:01	HLA-DRB4 Reference
HLA-DRB5	HLA00915	HLA-DRB5*01:01:01	HLA-DRB5 Reference

^aWhile each locus has at least one reference allele (e.g., HLA-A Reference), reference alleles for some allele families at a given locus are also identified (e.g., A*01 Reference). In some cases, the same allele may serve as a reference for multiple allele families.

Full gene sequence (i.e., for all exons, introns and UTRs) is available for all alleles on this table. Allele names for those alleles that include only two or three fields (e.g., HLA-B*73:01 or HLA-B*07:02:01) indicate that no synonymous or non-coding polymorphism, respectively, has been identified for those alleles as of Reference Database release version 3.25.0. Each allele family reference was selected on the basis of close sequence-identity between that reference allele and the alleles in that family.

Table 4

HLA Allele Remapping for WS Database LiffOver and Consensus Linking Functions

Reference Database Release in which the change occurred	Rationale	Original Allele Name	Original Accession Number	Current Allele Name	Current Accession Number	Current Allele Present in Version 3.25.0?	Accession Number Change?	WS Database Action When Original Allele Name Has Been Submitted
3.17.0	Sequence identical	B*49:15	HLA05834	B*49:01:01	HLA00340	YES	YES	
3.20.0	Sequence renamed	A*26:03:02	HLA04741	A*26:111	HLA04741	YES	NO	
3.21.0	Sequence error	DRB1*11:1:02	HLA02157	DRB1*11:1:01	HLA00765	YES	YES	
3.22.0	Sequence error	A*03:194	HLA11939	A*03:213	HLA12966	YES	YES	Use 3.25.0 version of Current Allele Name
3.24.0	Sequence error	A*23:69	HLA12676	A*23:01:01:01	HLA00048	YES ¹	YES	
	Sequence identical	DRB1*08:01:03	HLA02257	DRB1*08:01:01	HLA00723	YES	YES	
3.25.0	Sequence renamed	DRB1*04:94:02N	HLA14178	DRB1*04:212N	HLA14178	YES	NO	
	Sequence renamed	A*30:02:12	HLA09547	A*30:100	HLA14873	YES	YES	
3.26.0	Sequence error	C*17:01:01:01	HLA00481	C*17:01:01:02	HLA04311	YES	YES	Use 3.25.0 version of Original Allele Name
	Sequence renamed	DPB1*35:01:02	HLA04110	DPB1*62:01	HLA04110	NO ²	NO	
3.27.0	Sequence identical	DQB1*06:220	NA ³	DQB1*06:217	HLA16016	NO ²	NA ³	Change to DQB1*06:01:01 ⁴
	Sequence identical	DPA1*02:02:01	HLA00508	DPA1*02:07:01:01	HLA15619	NO ⁵	YES	Use 3.25.0 version of Original Allele Name
3.29.0	Sequence identical	DQB1*03:01:01:13	HLA07476	DQB1*03:01:01:07	HLA17167	NO ⁶	YES	Change to DQB1*03:01:01:01 ⁷

Changes to HLA allele names and their associated accession numbers that occurred in Reference Database versions 3.15.0 – 3.29.0, and the action taken by the WS Database when the original allele name is encountered are shown. The data in all but the last column are derived from the hla_nom.txt and alleleref_history.txt files in Reference Database version 3.28.0.

NA: Not applicable.

¹The allele name in Reference Database version 3.25.0 is A*23:01:01.

²This current allele name was assigned in Reference Database version 3.27.0.

³No accession number was released for the DQB1*06:220 allele; this allele name has not appeared in any release version.

⁴DQB1*06:01:01 is the lowest numbered allele sharing a common prefix (DQB1*06) with DQB1*06:220 (or DQB1*06:217).

⁵This current allele name was assigned in Reference Database version 3.28.0.

⁶Both the original and current allele names were assigned in Reference Database version 3.29.0.

DQB1*03:01:01:01 is the lowest numbered allele sharing a common prefix (DQB1*03:01:01:13 (or DQB1*03:01:01:07)

Author Manuscript

Author Manuscript

Author Manuscript

Author Manuscript