**Title:** Histoimmunogenetics Markup Language 1.0: Reporting Next Generation Sequencing-based HLA and KIR Genotyping

**Authors:** Robert P. Milius ¹, Michael Heuer ¹, Daniel Valiga ¹, Kathryn J. Doroschak ¹, Caleb J. Kennedy ¹, Yung Tsi Bolon ¹, Joel Schneider ¹, Jane Pollack ², Hwa Ran Kim ², Nezih Cereb ², Jill A. Hollenbach ³, Steven J. Mack ⁴, Martin Maiers ¹

1. National Marrow Donor Program, MN USA
2. HistoGenetics LLC, USA
3. University of California – San Francisco
4. Children's Hospital & Research Center Oakland, Oakland, CA, USA

**Communicating Author:**
Robert P. Milius
3001 Broadway St NE, Suite 100
National Marrow Donor Program
Minneapolis, MN 55413-1753 USA
(tel) +1-612-627-5884
(fax) +1-612-884-8677
bmilius@nmdp.org

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**Key Words:** HML; NGS; HLA; KIR; MIRING; data standards; genotyping

**Abbreviations:**
BRIDG: Biomedical Research Integrated Domain Group
CDISC: Clinical Data Interchange Standards Consortium
DaSH: Data Standard Hackathon
EMBL: European Molecular Biology Laboratory
ENA: European Nucleotide Archive
FDA: Food and Drug Administration
GL: Genotype List
HML: Histoimmunogenetics Markup Language
HLA: Human Leucocyte Antigen
IMGT: ImMunoGeneTics
ISO: International Organization for Standardization
LSDAM: Life Sciences Domain Analysis Model
KIR: Killer-cell Immunoglobulin-like Receptor
MHC: Major Histocompatibility Complex
MIRING: Minimum Information for Reporting Immunogenomic NGS Genotyping
NCI: National Cancer Institute
NGS: Next Generation Sequencing
NMDP: National Marrow Donor Program
OID: Object Identifier
SBT: Sequence Based Typing
SSO: Sequence Specific Oligonucleotide
SSP: Sequence Specific Primer
URI: Uniform Resource Identifier
XML: eXtensible Markup Language

**Abbreviated Title:** HML 1.0 For NGS HLA and KIR Genotyping
Abstract

We present an electronic format for exchanging data for HLA and KIR genotyping with extensions for next-generation sequencing (NGS). This format addresses NGS data exchange by refining the Histoimmunogenetics Markup Language (HML) to conform to the proposed Minimum Information for Reporting Immunogenomic NGS Genotyping (MIRING) reporting guidelines (miring.immunogenomics.org). Our refinements of HML include two major additions. First, NGS is supported by new XML structures to capture additional NGS data and metadata required to produce a genotyping result, including analysis-dependent (dynamic) and method-dependent (static) components. A full genotype, consensus sequence, and the surrounding metadata are included directly, while the raw sequence reads and platform documentation are externally referenced. Second, genotype ambiguity is fully represented by integrating Genotype List Strings, which use a hierarchical set of delimiters to represent allele and genotype ambiguity in a complete and accurate fashion. HML also continues to enable the transmission of legacy methods (e.g. site-specific oligonucleotide, sequence-specific priming, and sequence based typing (SBT)), adding features such as allowing multiple group-specific sequencing primers, and fully leveraging techniques that combine multiple methods to obtain a single result, such as SBT integrated with NGS.
1. Introduction

Human leukocyte antigen (HLA) genotyping is fundamental for research and clinical practice in immunogenetics and histocompatibility. Methods for generating this information have dramatically improved in the last three decades, from pioneering serological methods to modern DNA-based genotyping methods [1], revealing over time a seemingly never ending expansion of allele diversity [2], and presenting the subsequent challenge of reinterpretation of earlier results in light of new knowledge [3–6]. It has become increasingly clear that in order to ‘future-proof’ the results as much as possible, data standards are needed for recording and reporting of these results which include not only the allele assignments, but also the metadata (data describing other data) surrounding laboratory methods and allele assignment rules [5,7].

A new generation of sequencing methods has compounded this challenge. These methods are characterized by massively parallel technologies leading to high throughput sequencing, allowing for routine interrogation of not only select regions of a gene, but also whole-exon or whole-gene sequencing. While these technologies, commonly known as next-generation sequencing (NGS), have existed for several years, their application to HLA has been challenging due to a high degree of allelic polymorphism, the lack of robust genomic reference alignments for the MHC region, and the complex metadata required for allele assignment and later reinterpretation as reference sequences are refined and expanded. These considerations emphasize the need for recording and reporting complete metadata surrounding data collection, and in particular for data processing and interpretation.

Recently, a group of histocompatibility and immunogenetics stakeholders including clinicians, researchers, instrument manufacturers and software developers has gathered in a series of meetings to develop standards for recording and reporting NGS based genotyping of HLA [8]. One of the goals of these meetings has been to identify the Minimum Information for Reporting Immunogenomic NGS Genotyping (MIRING) [9] based on the principles of the reporting guidelines for the Minimum Information for Biological and Biomedical Investigations (MIBBI) [10]. The MIRING identifies eight principles for reporting NGS based genotyping of immunogenomics data. While the MIRING provides principles and guidelines, it does not provide a technical specification.

To develop a technical specification that meets these principles, we have extended the Histocompatibility Markup Language (HML) [11]. HML is an electronic messaging format based on XML and developed as a community standard allowing explicit representation of genotyping results for polymorphic immunogenetic gene systems such as HLA and KIR. Primary data in the form of Sequence Specific Oligonucleotide (SSO)/Sequence Specific Primer (SSP) reactivities and Sequence Based Typing (SBT) sequences can be linked with nomenclature-based allele assignment. In extending HML, we added the ability to include NGS based genotyping, including metadata surrounding the primary data and acquisition methods, consensus sequences, variant calling including novel polymorphisms, and reporting full allele and genotype ambiguity through the use of Genotype List (GL) Strings [12].

2. Methods

The development of HML 1.0 was led by the National Marrow Donor Program (NMDP) through a series of meetings and discussions with the HLA Information Exchange Data Format Standards Group, and later with the Immunogenomic NGS Data Consortium [8], a community of registries, clinical and research laboratories, and industry partners focused on identifying and addressing specific data-reporting requirements for NGS-based genotyping, and occurred in parallel with development of MIRING.

In September 2014, a community of stakeholders including researchers, clinicians, software developers, sequencing platform vendors, HLA and KIR sequence database developers and administrators, registry
and donor center IT professionals, and laboratory support staff gathered to attend a Data Standards Hackathon (DaSH) [13] to develop new ways to exchange NGS data for HLA and KIR. During this event the current state of HML was vetted by the community, and at the same time MIRING was further refined. These exercises led to new substantial requirements for the final specification for HML 1.0.

A rapid prototyping approach was used for the development of XML schemas for HML. As schemas were developed, example messages were created with various scenarios of HLA and KIR genotyping and syntactically validated using automated validation tools. The schema and examples can be found at https://bioinformatics.bethematchclinical.org/HLA-Resources/HML/.

In designing HML, it was important to accommodate the use of pointers to external references and datasets. This allows users to reuse previously registered metadata, and to reference datasets that are impractical to include in the HML message. These include registered methodologies, reference sequences, and raw sequence reads.

### 3. Results

HML 1.0 retains similar overall structure to previous versions, but with notable changes. As with earlier versions, reporting of primary data is separated from allele assignment. NGS is supported by new XML structures to capture all NGS data and metadata required to produce a genotyping result, including analysis-dependent (dynamic) and method-dependent (static) components. Pointers to external locations refer to registered methods, raw NGS reads, and reference standards. A separate component describing consensus sequences and variants was created specifically to accommodate NGS data, but could be used for other methodologies if desired. This component includes metadata describing the consensus sequences such as references sequences, phasing information, expected copy number, sequence block continuity, and other metadata. Reporting of allele assignment with full genotypic and allelic ambiguity is achieved through the use of GL Strings.

The overall HML 1.0 structure is seen in Figure 1. An HML message has four main sections: the document header, the typing methods, the allele assignment, and the consensus sequence. An HML message will contain the document metadata, and data describing one or more samples, generated with one or more typing methods for one or more gene families.

In addition to several minor improvements and many other components left to retain backward compatibility, two major improvements were introduced in HML 1.0. First, new elements that conform to the MIRING reporting guidelines were added. Second, GL strings were added to allow expression of full allelic and genotypic ambiguity in addition to, and eventually in lieu of, NMDP allele codes. Although the expectation is that NGS will lead to unambiguous allele assignment through full gene sequencing, earlier methodologies will continue to be used and reported in HML 1.0, and in the near term NGS based genotyping may be limited by some laboratories to interrogating specific exons leading to allele and genotype ambiguity. Additional changes include enhancements to Sanger-sequence based typing (SBT-Sanger), external references to typing kit information, and expansions to include multiple gene families (e.g., HLA, KIR).

In the specification below, note that the declared element and attribute names are in Courier font to highlight their abstract nature.

#### 3.1. HML Header

The header details are seen in Figure 2. Here, the message is contextualized, providing details such as identifying the lab sending the message, the report identifier, the sample(s) the report describes, and other reference information. This header does not contain details about any particular test, only the context in which it occurred.
3.1. hmlid

A unique identifier for the HML document can be included through the hmlid element. The hmlid follows the HL7 convention for instance identifiers using a two-part key [14]. An ISO Object Identifier (OID) [15] may be provided through the root attribute and often represents the unique organization identifier publicly registered for an organization. The extension attribute contains the unique document id managed internally by the reporting organization. Together, the root and extension attributes combine to provide a globally unique identifier.

Alternatively the ISO OID may be extended directly with the extension in the root, or a Universally Unique Identifier (UUID), also known as Globally Unique Identifier (GUID), may be provided in the root without an extension. Examples of these different ways of using hmlid are found in Figure 3.

3.1.2. reporting-center

The reporting-center element identifies the entity/organization sending the HML message. If included, it must contain a unique reporting-center-id identifying the sender. A new attribute, reporting-center-context, has been added. This is used to report the context/naming authority of the identifier. This can be used by different organizations to identify the reporting organization using registered contact information. For example, although this element is optional, NMDP requires the reporting-center element in an operational context. If the reporting-center-context is not included, it is assumed to be "NMDP" and is a unique identifier that is registered with the NMDP. Another example of the use of this element is to refer to a testing lab registered in the NCBI Genetic Testing Registry (GTR) [16] through the GTR Lab ID. Figure 4 shows examples of how reporting-center-id may be used.

3.1.3. typing-test-names

The typing-test-names element specifies a list of test names internally referenced by the SSO or SSP typing methods. It contains a test-id attribute associated with one or more typing-test-name containing test identifiers, which together are used to define a typing kit. It wraps a list of typing-test-name elements containing the test identifiers. The test-id is then used as an internal reference identifier to be used in the SSO and SSP typing methods sections.

3.1.4. sample

An HML message contains one or more sample elements enclosing the genotyping data pertaining to a particular sample. The id and center-code attributes identify the sample and the organization where the sample originated (e.g., transplant or donor center) respectively. Multiple samples from the same individual, or different individuals can be included in a single HML message. A collection-method element may be used to identify how the sample was collected, e.g., swab, filter paper, blood aliquot. Each sample may be associated with multiple typing elements.

3.1.4.1. typing

The typing element encapsulates the primary data from a genotyping method with a genotyping result (allele-assignment) that was determined from the primary data (typing-method) and/or consensus sequences (consensus-sequence). The gene-family attribute is used to constrain the typing to gene family name recognized by the HUGO Gene Nomenclature Committee [17] (e.g., HLA, KIR).

3.2. Genotyping Method and Primary Data

Figure 5 describes typing method and primary data elements. Here the method used to perform the genotyping and the primary un-interpreted results from that method are reported. This can be one of SSO, SSP, SBT-Sanger, or SBT-NGS.
3.2.1. Test IDs & Locus

To comply with MIRING requirements, three new attributes referencing additional metadata are available to more fully describe the method and locus targeted for each typing method. Two of these attributes name a test registrar (test-id-source) and an identifier (test-id) that can be dereferenced through the registrar. In previous versions of HML, sso and ssp elements referenced test information in the test-id element found in the header, which is used for encapsulating the test methods consisting of one or more individual tests or probe kits within the HML document. These new attributes allow labs to register their test method in an external testing registrars, such as the NCBI GTR [16].

The third new attribute for each typing method element is locus, which indicates the target of the typing method. Locus is an optional attribute. If present this must be a gene name recognized by the HUGO Gene Nomenclature Committee [17], e.g., HLA-A, HLA-DRB1, KIR2DL1.

3.2.2. sso & ssp

The sso and ssp elements describe SSO and SSP genotyping methods, respectively. These elements are unchanged from previous versions of HML except for the addition of test-id, test-id-source, and locus attributes (see above). For submissions to NMDP, a corresponding typing-test-names/typing-test-name structure found in the header is expected.

3.2.3. sbt-sanger

The sbt (Sequence-Based Typing) element found in previous versions of HML has been renamed to sbt-sanger. This reflects the recognition that both Sanger and NGS typing methods are sequence based, but with differences in the metadata needed to support them. Other changes include an additional child element to support sub-amplification primers and results. These primers are used to resolve ambiguities and may be used either concurrently with or after the amplification step. In addition, multiple GSSP elements (Group Specific Sequencing Primers), describing PCR primer sequences used to amplify polymorphic regions of sequences, are now allowed. In previous versions of HML, only a single GSSP was allowed.

3.2.4. sbt-ngs

The new sbt-ngs element describes Sequence Based Typing using high throughput methods, also known as NGS (next-generation sequencing). As with other methods, test-id, test-id-source, and locus attributes describe the method and the target locus of the test. Each sbt-ngs may contain multiple raw-reads child elements.

The raw-reads element references the raw sequence reads generated by an NGS platform. This data can be quite large even for relatively small regions of the genome. Because of this, if the raw-reads are provided, this information is not directly embedded in the HML document, but instead is referenced through a uri (Uniform Resource Identifier) attribute that points to an external location and a format attribute that describes how the data is stored (e.g., the NCBI Sequence Read Archive (SRA)). Other attributes include the availability of the raw reads (public, private, or with permission) and processing metadata indicating whether or not the data are pooled (pooled), if the adapters have been trimmed (adapterTrimmed), or whether low-quality sequences have been trimmed (qualityTrimmed).

3.3. Allele Assignment

Figure 6 describes the Allele Assignment section. Several changes from previous versions of HML were made here. The element that was previously named interpretation is now renamed allele-assignment to more accurately reflect its purpose. Also, this element includes new attributes that
describe the allele database or other source of the nomenclature (allele-db, e.g., IMGT-HLA Database) and release version of the database (allele-version, e.g., 3.19.0) that was used for the allele assignment.

Allele assignment may be reported in three different formats. As in previous versions of HML, haploid and genotype-list elements are available and unchanged. A new third element representing Genotype List (GL) Strings (glstring), which allows allele assignments with full genotype and allele ambiguity [12], has been added. The GL String may be directly included, or a URI resource identifier pointing to a service that provides the GL String (e.g., https://gl.nmdp.org) may be used. An example of how a GL String may be reported is seen in Figure 7.

3.4. Consensus Sequence

Figure 8 describes the Consensus Sequence section. This is a new structure in HML that was created to provide the result of an alignment or assembly of shorter sequence reads generated by a NGS platform. It is dependent on a number of dynamic factors such as the processing pipeline including software and inputs, and references sequences used for alignments. This is not considered to be primary data, but rather as a type of interpretation. For this reason, it is not included within the typing method section, but is considered separately. While this section was created with NGS in mind, it may be used by other sequence-based typing methods if desired.

This section is largely based on many of the data elements found in the Variant Call Format (VCF) [18] and metadata identified from the MIRING [9]. Note HML uses start and end attributes with a 0-based coordinate system with closed-open ranges, i.e., the range includes the start position, but excludes the end position. This convention is used in many genomic data models (e.g., Global Alliance for Genomics and Health schemas [19,20], UCSC browser [21], BED files) because of its programming benefits. In contrast, the VCF specification indexes the 1st base having position 1.

A sequence is reported directly through the sequence element. A pointer to an external reference may also be included, or as variant of a reference sequence. If a variant is reported, the structure provides a method to report variant effects. Position specific quality values may also be recorded.

3.4.1. reference-database & reference-sequence

Together, the reference-database and reference-sequence elements describe the database containing the reference sequence used within the consensus-sequence-blocks. The reference-database may point to whole genome builds (e.g., Genome Reference Consortium), allele databases (e.g., IMGT-HLA Database), or another database containing sequences that are used as a reference (e.g., GenBank, EMBL-ENA). Metadata surrounding the reference-database include the name, a description, the version, its availability, whether or not the database is curated, and a URI pointing to its external location.

For each reference-database, one or more reference-sequence elements may be described. An id is used to uniquely identify the reference-sequence and is referenced by the consensus-sequence-block. Other metadata that may be provided as attributes include the name of the sequence, the start and end positions, an accession id, and a URI pointing the sequence directly.

Examples of how reference-database and reference-sequence can be used are found in Figure 9.

3.4.2. consensus-sequence-block

A consensus-sequence-block encapsulates a DNA sequence consisting of IUPAC nucleotides [22] and variants against a reference sequence if an exact match is not found, and associated metadata. A
reference-sequence-id is required and points to a reference-sequence found within the HML document. Start and end positions identify a targeted region on the reference. Other optional metadata are provided through attributes that describe the strand being identified, a description of the target region (e.g., “HLA-A exon 3”), a phase set identifier to associate different consensus-sequence-blocks, a continuity flag to indicate whether a gap exists between blocks, and an expected-copy-number to indicated how many copies of the sequence block was expected.

A quality score for positions within the consensus sequence may be captured in the sequence-quality element, using start and end attributes indicating the region of interest. Quality metrics for sequence quality vary by platform reflecting differences in sequencing chemistry and digital processing. Additional metrics, such as coverage depth at the sequence- and variant-levels (see below), lack common standards and may be calculated differently depending on the analytical method chosen. For these reasons HML quality reporting does not impose strict specification criteria.

### 3.4.3. variant & variant-effect

A variant element must be included if the sequence does not completely match the reference-sequence. A typical use for this element is describing a novel allele. The available attributes for variant corresponds with many data elements used in VCF files [18] and in the MIRING. The id attribute is used to uniquely identify the variant within the HML document. The start and end identify the coordinates of the reference sequence where the variant occurs. The nucleotides found at this region of the reference and what they are replace with are identified with the reference-bases and alternate-bases attributes, respectively.

An alternative external reference for the variant may be reported using the uri attribute. This could be a location for a VCF file, or an alternative format that represents the same information.

A child of variant, the variant-effect element reports the effect of this variation. The effect or effects should be described using Sequence Ontology (SO) variant effect terms [23] (e.g., missense_variant, stop_gained, downstream_gene_variant). Additional attributes may be used to provide more information on the effect, e.g., severity, POLYPHEN prediction, SIFT score [24,25]. In addition to the term, a Human Genome Variation Society (HGVS) nomenclature string [26,27] may be included. A uri may be used to provide an external reference for this variant effect. An example of how variants may be captured is seen in Figure 10.

### 3.5. Extensibility

HML provides for extensibility using two methods. One is through the use of the property element that has been introduced as an optional child element for several nodes (hml, sample, typing, typing-method types, and allele-assignment). Here, name/value pairs (jointly defined and agreed upon by the creator and consumer of the HML document) may be entered. Additionally, property elements may contain custom nested structures of the creator's choosing to add additional information and details to the enclosing element. The other method of extending the HML specification is through the use of the anyAttribute type associated with the "consensus-sequence-block", "sequence", "variant", and "variant-effect" elements that allows additional attributes not specified by the schema.

### 4. Discussion

We present HML 1.0, an electronic format for exchanging data for genotyping of histoimmunogenetic markers such as HLA and KIR, with extensions for next-generation sequencing. These improvements equip the message with the mechanisms to collect new forms of typing data and accurately report genotype
and allele ambiguity, in a machine-readable format that tightly pairs the typing method and result for downstream analyses.

4.1. MIRING

We have extended and enhanced HML to implement the principles and guidelines outlined by the MIRING. Eight elements were identified that need to be addressed in an MIRING compliant genotyping report.

MIRING elements and corresponding example solutions in HML 1.0 are presented in Table 1. In several cases, the MIRING offers technical solutions for how specific elements may be addressed, but these translate into different solutions in HML. For example, the pipe-delimited descriptor found in MIRING is required for FASTA files. Often same information is found within the XML structure of HML and may not be explicitly needed. For example, in HML the consensus-sequence-blocks are captured sequentially, and each block contains within it all pertinent information within the attributes and child elements. Because of this, a separate identifier for the consensus sequence block to associate sequences with consensus sequences, references sequences, variants and associated metadata is not needed.

4.2. Validation

HML 1.0 has been finalized and the XML schema can be found on https://bioinformatics.bethematchclinical.org/HLA-Resources/HML/. Reporting of NGS based genotyping of HLA and KIR conforming to MIRING principles is possible using a combination of HML and semantic and syntactic validation tools, currently under development. A syntactically correct HML genotyping report may not pass organization specific business rules for acceptance or compliance to reporting guidelines such as MIRING. On the other hand, business-specific HML messages should conform to, and validate with, public XML schema definitions (XSDs). Transparent validation encourages a standards-based approach to mutual exchange of data for reproducible research and clinical application. Because of this, different validation tools will need to be developed depending on use cases and degree of semantic validation. For example, when a URI is presented to refer to external data and metadata, the validator will need to assess whether the link exists, and whether the information it points to makes sense and is interpretable. If HML is extended through the use of property elements and anyAttribute attributes, validation is further complicated since they are largely ignored with syntactic validation tools. Robust validation tools for these special use cases must then be created. A tool to validate whether a given instance of a HML message is MIRING compliant is being developed.

4.3. Future work

HML 1.0 has been developed to serve the needs of the immunogenomics community. However we recognize the need for interoperability with the larger healthcare community and the potential to interface with EMR systems. In light of this, the possibility to encapsulate HML in HL7 messages or clinical structured documents is being explored.

HML is a standard derived from the community members listed above. However, the NMDP has a commitment to using standards developed and maintained by other parties whenever possible. HL7 is an international standards development organization committed to the interchange of healthcare data. We are investigating the utility of OIDs (referred to above) as unique identifiers, as well as HL7 interchange formats such as Clinical Document Architecture (CDA) and Fast Healthcare Interoperability Resources (FHIR) to encapsulate HML. There is an active community to develop these standards that serves a larger audience than the parties listed in this document. We are actively engaged with the HL7 Clinical Genomics Working Group to ensure the content contained in the HML 1.0 specification can be contained in an HL7 message. This will allow the parties listed here to leverage the standards developed and implemented by the HL7 community. HL7 is also a party to the Biomedical Research Integrated Domain
Group (BRIDG) as a consortium member including the FDA, CDISC, and NCI. The BRIDG members are developing artifacts to assist in the interchange of study-driven protocols and their regulatory artifacts, but also to assist in the interchange of molecular testing as part of the LSDAM (Life Sciences Domain Analysis Model). Since the HML 1.0 model is, at its core, an encapsulation of the results of a molecular test, we wish to leverage the work of those parties as well.

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6. References


HL7 Implementation Guidance for Unique Object Identifiers (OIDs), Release 1 2009:1–34.  


7. **Figure Legends**

- **Figure 1:** HML overall structure
- **Figure 2:** Header
- **Figure 3:** Examples of how hmlid can be used
- **Figure 4:** Examples of how reporting-center can be used
- **Figure 5:** Typing methods & primary data
- **Figure 6:** Allele Assignment
- **Figure 7:** Example of how allele-assignment may be used with glstring
- **Figure 8:** Consensus sequence
- **Figure 9:** Examples of using reference-database and reference-sequence
- **Figure 10:** Example of consensus-sequence-block with variants
8. Figures

Fig 1
Fig 2

```
  html
  version *
  project-name

  [ 0..1 ]
  hmlid
  root *
  extension

  [ 0..1 ]
  reporting-center
  reporting-center-id *
  reporting-center-context

  [ 1..* ]
  sample
  id *
  center-code

  [ 0..* ]
  typing-test-names
  test-id *

  [ 0..* ]
  collection-method

  [ 1..* ]
  typing
  gene-family *
  date *

  [ 1 ]
  Typing Method

  [ 0..* ]
  Allele Assignment

  [ 0..* ]
  Consensus Sequence
```

```
element
required attribute *
optional attribute
```
**OID using separate root and extension**
<hmlid extension="123456789" root="2.16.840.1.113883.3.1470"/>

**OID using only root**
<hmlid root="2.16.840.1.113883.3.1470.123456789"/>

**UUID/GUID**
<hmlid root="1ee83ff1-08ab-4fe7-b573-ea777e9bad51"/>
**NMDP defined reporting center without explicit context**
<reporting-center
  reporting-center-id="888" />

**NMDP defined reporting center with explicit context**
<reporting-center
  reporting-center-id="888"
  reporting-center-context="NMDP"/>

**Lab ID registered with the NCBI Genetic Testing Registry**
<reporting-center
  reporting-center-id="000000"
  reporting-center-context="NCBI GTR"/>
Fig 7

<allele-assignment
date="2015-01-26"
allele-db="IMGT/HLA"
allele-version="3.19.0">
<glstring>
HLA-A*01:01:01+HLA-A*24:02:01
</glstring>
</allele-assignment>
### Genome Reference Consortium

```
<reference-database
  name="GRCh38.p1"
  description="Genome Reference Consortium (GRC)"
  version="GRCh38.p1"
  availability="public"
  curated="true"
  <reference-sequence
    id="1"
    name="HSCHR6_MHC_MCF_CTG1"
    start="0"
    end="4827813"
    accession="GL000254.2"
  <reference-sequence
    id="2"/>
</reference-database>
```

### IMGT/HLA

```
<reference-database
  name="imgt-hla"
  description="IMGT/HLA Database"
  version="3.19.0"
  availability="public"
  curated="true"
  uri="http://www.ebi.ac.uk/ipd/imgt/hla">
  <reference-sequence
    id="3"
    name="HLA-A*01:01:01:01"
    start="0"
    end="3053"
    accession="HLA00001"
    uri="http://www.ebi.ac.uk/Tools/dbfetch/dbfetch?db=imgthla;id=HLA00001"/>
  <reference-sequence
    id="4"/>
</reference-database>
```

### No Database Reference

```
<reference-database availability="none">
  <reference-sequence id="0"/>
</reference-database>
```
<consensus-sequence-block
  reference-sequence-id="1"
  start="29942412" end="29945809"
  strand="1" phase-set="1"
  continuity="true" expected-copy-number="1"
  description="HLA-A full gene">
  <sequence> TTCCTGGATACTCACG ... etc </sequence>
  <!-- insertion -->
  <variant
    reference-bases="A"
    alternate-bases="AT"
    start="29945809" end="29945810">
    <variant-effect term="stop_gained"/>
  </variant>
  <!-- deletion -->
  <variant
    reference-bases="GG"
    alternate-bases="G"
    start="29942940" end="29942942">
    <variant-effect term="frameshift_variant"/>
  </variant>
  <sequence-quality
    sequence-start="29942412"
    sequence-end="29942612"
    quality-score="0.9"/>
  <sequence-quality
    sequence-start="29942612"
    sequence-end="29942942"
    quality-score="1.0"/>
</consensus-sequence-block>
## 9. Tables

### Table 1: Mapping of MIRING elements to HML solutions

<table>
<thead>
<tr>
<th>MIRING Elements</th>
<th>HML Solution Example with comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Message Annotation</td>
<td></td>
</tr>
<tr>
<td>1.1. Unique MIRING Message Identifier</td>
<td>&lt;hmlid root=&quot;x.x.x&quot; extension=&quot;x&quot; /&gt;</td>
</tr>
<tr>
<td></td>
<td>root may be ISO Object Identifier (OID) or Universally Unique Identifier (UUID)</td>
</tr>
<tr>
<td>1.2. Message Generator Contact Information</td>
<td>Using an NCBI GTR Lab ID:</td>
</tr>
<tr>
<td></td>
<td>&lt;reporting-center reporting-center-id=&quot;000000&quot; reporting-center-context=&quot;NCBI-GTR&quot; /&gt;</td>
</tr>
<tr>
<td>1.3. Platform Documentation</td>
<td>Using an NCBI GTR Test ID:</td>
</tr>
<tr>
<td></td>
<td>&lt;sbt-ngs ... test-id=&quot;GTR000000000.0&quot; test-id-source=&quot;NCBI-GTR&quot; /&gt;</td>
</tr>
<tr>
<td>1.4. Read Processing Documentation Reference</td>
<td>Using an NCBI GTR Test ID:</td>
</tr>
<tr>
<td></td>
<td>&lt;sbt-ngs ... test-id=&quot;GTR000000000.0&quot; test-id-source=&quot;NCBI-GTR&quot; /&gt;</td>
</tr>
<tr>
<td>1.5. Primary Data Availability</td>
<td>&lt;raw-reads ... availability=&quot;true&quot; ... /&gt;</td>
</tr>
<tr>
<td>1.6. Primary Data Reference</td>
<td>&lt;raw-reads ... uri=&quot;<a href="http://www.ncbi.nlm.nih.gov/sra/SRX0000000">http://www.ncbi.nlm.nih.gov/sra/SRX0000000</a>&quot; ... /&gt;</td>
</tr>
<tr>
<td>2. Reference Context</td>
<td></td>
</tr>
<tr>
<td>2.1. Reference Sequence Database Version for Allele Calling</td>
<td>For allele assignment, the allele database and version is captured in attributes:</td>
</tr>
<tr>
<td></td>
<td>&lt;allele-assignment ... allele-db=&quot;IMGT-HLADB&quot; allele-version=&quot;3.19.0&quot; &gt; &lt;glstring&gt;...&lt;/glstring&gt; &lt;/allele-assignment&gt;</td>
</tr>
<tr>
<td></td>
<td>For sequence variant, the reference database and sequence is identified through the reference-database and reference-sequence elements:</td>
</tr>
<tr>
<td></td>
<td>&lt;reference-database&gt; &lt;reference-sequence ... /&gt; &lt;reference-sequence ... /&gt; &lt;/reference-database&gt;</td>
</tr>
<tr>
<td>2.2. Individual Reference Sequences Applied</td>
<td>&lt;reference-sequence id=&quot;0&quot; name=&quot;HSCHR6_MHC_MCF_CTG1&quot; start=&quot;0&quot; end=&quot;4827813&quot; accession=&quot;CCDS34373.12&quot; /&gt;</td>
</tr>
<tr>
<td>Section</td>
<td>Description</td>
</tr>
<tr>
<td>---------</td>
<td>-------------</td>
</tr>
<tr>
<td>2.2.1</td>
<td>Reference Sequence Identifier</td>
</tr>
<tr>
<td>2.3</td>
<td>Reference Sequence Source Type</td>
</tr>
<tr>
<td>3</td>
<td>Full Genotype</td>
</tr>
<tr>
<td>3.1</td>
<td>Pertinent Locus/Loci</td>
</tr>
<tr>
<td>3.2</td>
<td>Formatted Genotype</td>
</tr>
<tr>
<td>3.3</td>
<td>Genotype Uniform Resource Identifier (URI)</td>
</tr>
<tr>
<td>4</td>
<td>Consensus Sequence</td>
</tr>
<tr>
<td>4.1</td>
<td></td>
</tr>
<tr>
<td>4.2</td>
<td>Consensus Sequence Descriptor</td>
</tr>
<tr>
<td>4.2.1</td>
<td>Consensus Sequence Block Identifier</td>
</tr>
<tr>
<td>4.2.2</td>
<td>Reference Sequence Identifier</td>
</tr>
<tr>
<td>4.2.3</td>
<td>Reference Sequence Coordinate</td>
</tr>
<tr>
<td>4.2.4</td>
<td>Phase Set</td>
</tr>
</tbody>
</table>
### 4.2.5 Copy Number

<table>
<thead>
<tr>
<th>Tag</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;consensus-sequence-block expected-copy-number=&quot;1&quot; ...</td>
<td>... &gt; ... &lt;/consensus-sequence-block&gt;</td>
</tr>
</tbody>
</table>

#### Reference Sequence Match

- **Reference sequence match = true**
  
  \[
  \begin{align*}
  \text{<consensus-sequence-block ... > } \\
  \text{<sequence> ... </sequence>} \\
  \text{</consensus-sequence-block>}
  \end{align*}
  \]

- **Reference sequence match = false**
  
  \[
  \begin{align*}
  \text{<consensus-sequence-block ... > } \\
  \text{<sequence> ... </sequence>} \\
  \text{<variant ... > ... </variant>} \\
  \text{</consensus-sequence-block>}
  \end{align*}
  \]

### 4.2.6 Sequence Continuity

<table>
<thead>
<tr>
<th>Tag</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;consensus-sequence-block ... continuity=&quot;true&quot; ...</td>
<td>... &gt;</td>
</tr>
</tbody>
</table>

### 5 Novel Polymorphisms

#### Reference

For allele assignment, the allele database and version is captured in attributes:

\[
\begin{align*}
\text{<allele-assignment ... } \\
\text{allele-db="IMGT-HLADB" } \\
\text{allele-version="3.19.0" } \\
\text{<glstring>...</glstring>} \\
\text{</allele-assignment>}
\end{align*}
\]

For sequence variant, the reference sequence is identified through the reference-database and reference-sequence elements.

\[
\begin{align*}
\text{<reference-database> } \\
\text{<reference-sequence ... >/} \\
\text{<reference-sequence ... >/} \\
\text{</reference-database>}
\end{align*}
\]

#### 5.1 Position

<table>
<thead>
<tr>
<th>Tag</th>
<th>Content</th>
</tr>
</thead>
</table>
| <variant start="123" end="124" ... | ... />

#### 5.3 Variant Identifier

<table>
<thead>
<tr>
<th>Tag</th>
<th>Content</th>
</tr>
</thead>
</table>
| <variant id="0" ... | ... />

#### 5.4 Reference Sequence

<table>
<thead>
<tr>
<th>Tag</th>
<th>Content</th>
</tr>
</thead>
</table>
| <variant reference-bases="A" ... | ... />

#### 5.5 Variant Sequence

<table>
<thead>
<tr>
<th>Tag</th>
<th>Content</th>
</tr>
</thead>
</table>
| <variant alternate-bases="TCG" ... | ... />

#### 5.6 Quality Score

<table>
<thead>
<tr>
<th>Tag</th>
<th>Content</th>
</tr>
</thead>
</table>
| <variant quality="90" | ... />

#### 5.7 Quality Filter Status

<table>
<thead>
<tr>
<th>Tag</th>
<th>Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;variant filter=&quot;pass&quot;</td>
<td>...</td>
</tr>
</tbody>
</table>

#### 5.8 INSDC Accession Number

While the accession number for the reference sequence can be provided in HML, there is no provision for providing one for the variant sequence. If an accession number is available, then for the purposes of the report, it is considered a reference and matched. Since this is optional ("when possible") in the MIRING, we do not see this as an issue. If a separate accession number for the variant is needed, consensus-sequence-block may be extended through the anyAttribute attribute, but this must be agreed upon by the creator and consumer of the HML message.

#### 5.9 Platform Documentation

Using an NCBI GTR Test ID:

\[
\begin{align*}
\text{<sbt-NGS ... } \\
\text{test-id="GTR000000000.0"}
\end{align*}
\]
<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
</table>
| **Read Processing Documentation** | NCBI GTR should be used for describing overall methods and pipeline strategy, i.e., standard methods and SOPs for running the test.  
<sbt-ngs _  
test-id="GTR000000000.0"  
test-id-source="NCBI-GTR" />  
Dynamic methodology that can change with each sequencing run should be captured with an Sequence Read Archive like data structure that is reported through the raw-reads uri attribute:  
<raw-reads _  
| **Primary Data** | <raw-reads _  