Minimum Information for Reporting Immunogenomic NGS Genotyping

Define an NGS Genotyping Result with 8 Categories of Information

MIRING Annotation
Reference Context
Full Genotype
Consensus Sequence
Novel Polymorphisms
Platform Documentation
Read Processing Documentation
Primary Data

MIRING Message
Accessory Data

http://igdawg.org/miring.html
MIRING & HML

- MIRING is a set of guiding principles

- HML 1.0 is a technical specification that allows information conforming to MIRING principles to be exchanged
  - But HML doesn’t have to be MIRING compliant
  - You can perfectly valid HML, yet doesn’t contain much information.
  - XML Schema
An **XML Schema** is a language for expressing constraints about **XML** documents. There are several different schema languages in widespread use, but the main ones are Document Type Definitions (DTDs), Relax-NG, Schematron and W3C **XSD** (**XML Schema Definitions**).

**XML Schema - W3C**

[https://www.w3.org/standards/xml/schema](https://www.w3.org/standards/xml/schema)

World Wide Web Consortium
Histoimmunogenetics Markup Language 1.0: Reporting next generation sequencing-based HLA and KIR genotyping

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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.

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For more on HML, see the Bioinformatics HML information page.

Public XML Schemas

Schemas on this page can be referenced either using the "http" OR "https" protocols. Some software may be unable to negotiate a secure connection if using "https" so be aware of your audience. All submissions of HML data should be sent over a secure transport layer.

- HML 0.2 (diagram) **
- HML 0.3 (diagram) **
- HML 0.3.3 (diagram) **
- HML 1.0 (diagram)
- **HML 1.0.1 (diagram) **
Histoimmunogenetics Markup Language (HML)
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> http://www.gnu.org/licenses/lgpl.html

-->

```xml
<xs:schema xmlns="http://schemas.nmdp.org/spec/hml/1.0.1"
    xmlns:hml="http://schemas.nmdp.org/spec/hml/1.0.1" xmlns:xs="http://www.w3.org/2001/XMLSchema"
    elementFormDefault="qualified" attributeFormDefault="unqualified"
    targetNamespace="http://schemas.nmdp.org/spec/hml/1.0.1" version="1.0.1">
  <!-- HML document root -->
  <xs:element name="hml" id="hml" nillable="false">
    <xs:annotation>
      <xs:documentation>
        Root element of the document identifying it as an HML message. Must contain the version of HML that the modeled data in this document uses. Children: --------- - property (optional, qty: 0 or more) - hmlid (optional, qty: 1) - reporting-center (optional, qty: 1) - Required for NMDP samples - sample (required, qty: 1 or more) - typing-test-names (optional, qty: 0 or more) Attributes: --------- - version: Version of HML the document follows (required) - project-name: Name of the typing project (optional)
      </xs:documentation>
    </xs:annotation>
  </xs:element>
</xs:schema>
```

<xs:complexType>
  <xs:sequence>
    <xs:element name="property" type="hmlns:property" minOccurs="0" maxOccurs="unbounded"/>
    <xs:element name="hmlid" type="hmlns:hmlid" minOccurs="0" maxOccurs="1"/>
    <xs:element ref="reporting-center" minOccurs="0" maxOccurs="1"/>
  </xs:sequence>
</xs:complexType>
```
For more on HML, see the Bioinformatics HML information page.

Public XML Schemas

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- HML 0.2 (diagram)**
- HML 0.3 (diagram)**
- HML 0.3.3 (diagram)**
- HML 1.0 (diagram)
- HML 1.0.1 (diagram)**
Many elements are not required to be valid HML
Tiny Valid HML 1.0.1

```xml
<?xml version="1.0" ?>
<hml xmlns="http://schemas.nmdp.org/spec/hml/1.0.1"
    version="1.0.1">
    <sample id="1">
        <typing gene-family="HLA" date="2016-07-01">
            <typing-method/>
        </typing>
    </sample>
</hml>
```
Command line validation

$ xmlint --noout --schema hml-1.0.1.xsd tiny.hml

tiny.hml validates

$
MIRING Validator

- First, is it valid HML?
  - Is it well formed?
  - XML Schema

- Second, does it contain elements necessary to support MIRING?
  - Schematron
An **XML Schema** is a language for expressing constraints about XML documents. There are several different schema languages in widespread use, but the main ones are Document Type Definitions (DTDs), Relax-NG, **Schematron** and W3C **XSD** (**XML Schema** Definitions).

**XML Schema - W3C**

https://www.w3.org/standards/xml/schema

World Wide Web Consortium
XML Schema vs Schematron

• XML Schema
  – Grammar based
    • e.g., cardinality, data types, value sets
  – XSD
  – W3C

• Schematron
  – Rules based on patterns
    • Assertions applied to a specific context within the document
    • e.g., “start” must be smaller than “end”
  – XSLT and XPath
  – ISO/IEC
Tiers of Validation

- **Tier 1**
  - XML Schema to validate both MIRING and HML schemas. We can check simple MIRING and HML syntax rules, for existence of nodes or attributes.

- **Tier 2**
  - Use Schematron to validate some more complicated rules. More complicated data comparisons (Start < End) can be done here. Schematron is an XML validation rules engine, so we use it to do comparisons, etc. that XML Schema validation can't do.

- **Tier 3**
  - Tier 3 is where we assigned external lookups, such as Method Registries, Reference Sequence Databases, GLString validations, etc. This is outside of scope for now.
MIRING Validator for HML

Version 1.1.5

Paste some XML, compliant with HML Version that corresponds to your HML into the HML field and hit [ VALIDATE ].

Or, use the [ BROWSE ] button find an HML document on your computer.

A MIRING Results Report will appear below.

HML Input

```xml
<?xml version="1.0" ?>
<html xmlns="http://schemas.nmdp.org/spec/hml/1.0.1" version="1.0.1">
  <sample id="123">
    <typing gene-family="HLA" date="2016-07-01">
      <typing-method/>
    </typing>
  </sample>
</html>
```

MIRING Results

✔ Passed XML Schema Validation for HML 1.0

⚠ NOT MIRING Compliant
MIRING Results

✔ Passed XML Schema Validation for HML 1.0

⚠ NOT MIRING Compliant

Use the MIRING Validator in your own application:

The MIRING HML Validator is a REST service, available to the public, hosted at miring.b12x.org/validator/ValidateMiring/

You can use this service programmatically using a POST request, with a single form parameter, "xml!", containing the HML text.

Try this example using cURL:

```
$ curl -X POST --data-urlencode 'xml[]=<hml>...</hml>' http://miring.b12x.org/validator/ValidateMiring/
```

Validator source is on GITHUB

More information on MIRING, HML, and validation
HOW TO READ A VALIDATION LINE

Example of a validation line

```html
hml-result severity="fatal"
```

description[306,138] Value '' with length = '0' is not facet-valid with respect to minLength '5' for type '
#AnonType_uriraw-reads'. /description

solution Verify that your HML file is well formed, and conforms to
http://schemas.nmdp.org/spec/hml/1.0.1/hml-1.0.1.xsd /solution /hml-result

```html
hml-result severity="fatal"
```

This line shows the hml or miring result which is split into two sections. The severity levels allows for the validator to know if the issue should be allowed or not this can either be fatal, hml, miring, warning, or info. HML, miring, and fatal errors will stop the validation and must be fixed immediately. For miring results, you will see a miring-rule-id which specifies which rule is being violated.

[306,138] Represents the row and column number respectfully (Note: If you are missing a matching end tag the system might report the location to be the end of the file. Please just refer to the description if this happens)

Value '' with length = '0' is not facet-valid with respect to minLength '5' for type '
#AnonType_uriraw-reads'. is an autogenerated error message based from SAXParser Schema Error codes this will always appear in the description nodes

Verify that your HML file is well formed, and conforms to http://schemas.nmdp.org/spec/hml/1.0.1/hml-
1.0.1.xsd is a statement made by the developers on a start on how to fix this specific issue and will always be in the solution node

On occasions you will see a location, this will be an XPath leading to the error, I suggest reading this webpage on how to read it
# Rules

<table>
<thead>
<tr>
<th>Miring Element</th>
<th>Miring Description</th>
<th>Rule ID</th>
<th>Severity</th>
<th>Rule Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Message Annotation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.1</td>
<td>Unique Miring Message Identifier</td>
<td>1.1.a</td>
<td>miring</td>
<td>Document has exactly one hmlid node directly under the root hml node</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1.1.c</td>
<td>info</td>
<td>if hmlid is an OID the hmlid node should have a root and an extension attribute.</td>
</tr>
<tr>
<td>1.2</td>
<td>Message Generator Contact</td>
<td>1.2.a</td>
<td>miring</td>
<td>Document has exactly one reporting-center node directly under the root hml node</td>
</tr>
<tr>
<td></td>
<td>Information</td>
<td>1.2.b</td>
<td>warning</td>
<td>reporting-center has one reporting-center-id and one reporting-center-context attributes</td>
</tr>
<tr>
<td>1.3</td>
<td>Platform Documentation (MIRING</td>
<td>1.3.a</td>
<td>miring</td>
<td>Every sso ssp sbt-sanger sbt-ngs node should have a test-id and test-id-source attributes</td>
</tr>
<tr>
<td></td>
<td>element 6) Reference</td>
<td>1.3.b</td>
<td>info</td>
<td>test-id and test-id-source are in a recognized format. NCBI-GTR</td>
</tr>
<tr>
<td>1.4</td>
<td>Read Processing Documentation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MIRING elements</td>
<td>HML solution example with comment</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>----------------</td>
<td>----------------------------------</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
| 1 Message Annotation | <hmlid root="x.x.x" extension="x" />
| 1.1 Unique MIRING Message Identifier | root may be ISO Object Identifier (OID) or Universally Unique Identifier (UUID) |
| 1.2 Message Generator Contact Information | Using an NCBI GTR Lab ID:
| | <reporting-center
| | reporting-center-id="000000"
| | reporting-center-context="NCBI-GTR" />
| 1.3 Platform Documentation | Using an NCBI GTR Test ID:
| | <sbt-ngs...
| | test-id="GTR000000000.0"
| | test-id-source="NCBI-GTR" />
| 1.4 Read Processing Documentation Reference | Using an NCBI GTR Test ID:
| | <sbt-ngs...
| | test-id="GTR000000000.0"
| | test-id-source="NCBI-GTR" />
| 1.5 Primary Data Availability | <raw-reads ... availability="true".../>
| 1.6 Primary Data Reference | <raw-reads...
| 2 Reference Context | |
| 2.1 Reference Sequence Database Version for Allele Calling | For allele assignment, the allele database and version is captured in attributes:
| | <allele-assignment...
| | allele-db="TMGT-HLAAB"
| | allele-version="5.19.0">
| | <glistings...
| | glistings>
| | </allele-assignment>
| | For sequence variant, the reference database and sequence is identified through the reference-database and reference-sequence elements.
| | <reference-database>
| | <reference-sequence>...
| | </reference-database> |
| 2.2 Individual Reference Sequences Applied | <reference-sequence id="0"
| | name="HSCHR6_HHC_MCP_OTG1"
| | start="0"
| | end="4827813"
| | accession="CCDS34373.18"
| 2.2.1 Reference Sequence Identifier | <reference-sequence ... id="0".../> |
| 2.3 Reference Sequence Source Type | <reference-database
| | public="true"
| | curated="true"/> |
Things to remember…

- Just because your HML passes both HML and MIRING validation doesn’t mean it’s okay

- The validator doesn’t check…
  - if your coordinates are 0-base
  - if your reference URIs and IDs match with the coordinates you supply
  - if the URIs you enter actually resolve to anything real
  - if the strand you enter is correct for the reference you supply
  - If the variant coordinates are within the window of the reference sequence that makes up your consensus-sequence-block
  - etc

- The creator of the HML is still responsible for its content, that it’s complete and makes sense
Links

• Web GUI
  – http://miring.b12x.org

• Source code
  – https://github.com/nmdp-bioinformatics/MiringValidator

• HML manuscript

• MIRING manuscript
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  – Chia-Jung Chang
  – Marcelo Fernandez-Vina

• (please let know who else…)

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Thank you!

Questions?