airr-standards Documentation

Release 1.3

AIRR Community

Oct 13, 2020

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The Adaptive Immune Receptor Repertoire (AIRR) Community of The Antibody Society is a research-driven group that is organizing and coordinating stakeholders in the use of next-generation sequencing (NGS) technologies to study antibody/B-cell and T-cell receptor repertoires. Recent advances in sequencing technology have made it possible to sample the immune repertoire in exquisite detail. AIRR sequencing has enormous promise for understanding the dynamics of the immune repertoire in vaccinology, infectious disease, autoimmunity, and cancer biology, but also poses substantial challenges. The AIRR Community was established to meet these challenges.

CHAPTER 1

Introduction to the AIRR Standards

The AIRR Community is developing a set of standards for describing, reporting, storing, and sharing adaptive immune receptor repertoire (AIRR) data, such as sequences of antibodies and T cell receptors (TCRs). Some specific efforts include:

- The MiAIRR standard for describing minimal information about AIRR datasets, including sample collection and data processing information.
- Data submission guidelines and workflows.
- Data representations (file format) specifications for storing large amounts of annotated AIRR data.
- API to query and download AIRR data from repositories/databases as part of the AIRR Data Commons.
- A community standard for software tools which will allow conforming tools to gain community recognition.
- Set of reference software tools for reading, writing and validating data in the AIRR standards.
- · A database and web submission frontend for inferred germline genes

CHAPTER 2

Table of Contents

2.1 Getting Started

This website provides information and resources regarding the AIRR Community Standards for the diverse community of immunology researchers, bioinformaticians, and software developers.

2.1.1 MiAIRR standard for study data submission

- Gather experimental and analysis information about your study to conform to the *MiAIRR* standard (minimal information about adaptive immune receptor repertoires).
- Submission of your study data to a public repository.

2.1.2 AIRR Data Commons for query and download of AIRR-seq data

• Query publicly available AIRR-seq studies in the AIRR Data Commons.

2.1.3 Resources related to data representations and software development

- Schema, definitions and file formats for the *AIRR Data Model*. The AIRR Data Model defines the structure and relationship for the MiAIRR data elements.
- Software guidelines for tools developers to enable rigorous and reproducible immune repertoire research.
- AIRR Data Commons API provides programmatic access to query and download AIRR-seq data.

2.1.4 Software tools and libraries

- Python reference library for reading/writing/validating AIRR data files.
- R reference library for reading/writing/validating AIRR data files.

- ADC API reference implementation for a local data repository.
- Resources and tools that support the AIRR Standards.

2.1.5 Tutorials, examples and workflows

AIRR Rearrangement TSV Interoperability Example

The example that follows illustrates the interoperability provided by the AIRR Rearrangement schema. The code provided demonstrates how to take AIRR formatted data output by IgBLAST and combine it with data processed by IMGT/HighV-QUEST that has converted to the AIRR format by Change-O. Then, the merged output of these two distinct tools is used to (a) create MiAIRR compliant GenBank/TLS submission files, and (b) perform a simple V gene usage analysis task.

Data

We've hosted a small set of example data from BioProject PRJNA338795 (Vander Heiden et al, 2017. J Immunol.) containing both input and output of the example. It may be downloaded from:

Example Data

Walkthrough

Environment setup

We'll use the Immcantation docker image for this example, which comes loaded with all the tools used in the steps that follow:

```
# Download the image
docker pull kleinstein/immcantation:devel
# Invoke a shell session inside the Immcantation docker image
# Map example data (~/data) to the container's /data directory
$> docker run -it -v ~/data:/data:z kleinstein/immcantation:devel bash
```

Generate AIRR formatted TSV files

TSV files compliant with the AIRR Rearrangement schema may be output directly from IgBLAST v1.9+ or generated from IMGT/HighV-QUEST output (or IgBLAST <=1.8 ouput) using the MakeDb parser provided by Change-O:

```
# Generate TSV directly with IgBLAST
$> cd /data
$> export IGDATA=/usr/local/share/igblast
$> igblastn -query HD13M.fasta -out HD13M_fmt19.tsv -outfmt 19 \
    -germline_db_V $IGDATA/database/imgt_human_ig_v \
    -germline_db_D $IGDATA/database/imgt_human_ig_d \
    -germline_db_J $IGDATA/database/imgt_human_ig_j \
    -auxiliary_data $IGDATA/optional_file/human_gl.aux \
    -ig_seqtype Ig -organism human \
    -domain_system imgt
```

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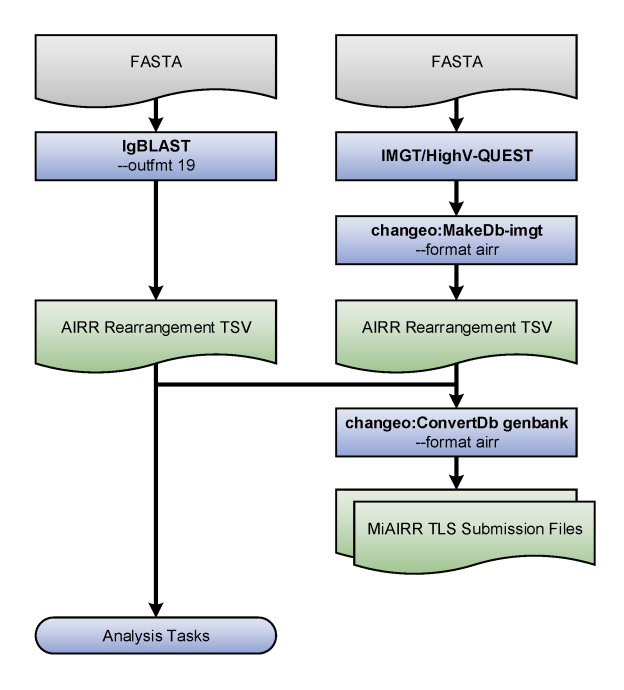


Fig. 1: Flowchart of the example steps.

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Generate GenBank/TLS submission files

AIRR TSV files can be input directly in Change-O's ConvertDb-genbank tool to generate MiAIRR compliant files for submission to GenBank/TLS:

Merge files and count V family usage

AIRR TSV files from different tools and easy combined to perform analysis on data generated using different software. Below is shown a simple V family usage analysis after merging the IgBLAST and IMGT/HighV-QUEST outputs into a single table:

```
# Count V family usage in R
# Imports
$> R
R> library(alakazam)
R> library(dplyr)
R> library(ggplot2)
# Merge IgBLAST and IMGT/HighV-QUEST results
R> db_m <- read.delim("HD13M_fmt7_db-pass.tsv")</pre>
R> db_n <- read.delim("HD13N_imgt_db-pass.tsv")</pre>
R> db_m$cell_type <- "memory"</pre>
R> db_n$cell_type <- "naive"</pre>
R> db <- bind_rows(db_m, db_n)</pre>
# Subset to heavy chain
R> db <- subset(db, grepl("IGH", v_call))</pre>
# Count combined V gene usage
R> v_usage <- countGenes(db, "v_call", groups="cell_type",</pre>
                           mode="family")
# Plot V family usage
R> ggplot(v_usage, aes(x=GENE, y=SEQ_FREQ, fill=cell_type)) +
    geom_col(position="dodge") +
    scale_fill_brewer(name="Cell type", palette="Set1") +
```

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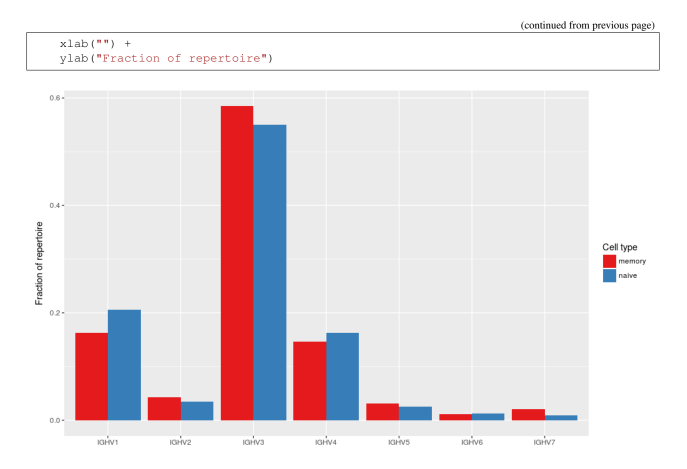


Fig. 2: V family usage for the combined data set.

ADC API Query and Analysis Example

This example shows how repertoires and associated rearrangments may be queried from a data repository using the ADC API and then a simple analysis is performed. The example is split between two python scripts; one that performs the query and saves the data into files, and another that reads the data from the files and generates a grouped CDR3 amino acid length distribution plot. The two scripts could be combined into one, but this example illustrates how the data can be saved into files for later use. The example uses the AIRR standards python library.

Data

This example retrieves data for the following study, which is identified by NCBI BioProject PRJNA300878. In this example, we are only going to query and retrieve the T cell repertoires.

Rubelt, F. et al., 2016. Individual heritable differences result in unique cell lymphocyte receptor repertoires of naive and antigen-experienced cells. Nature communications, 7, p.11112.

Basic study description:

- 5 pairs of human twins
- B-cells and T-cells sequenced
- · B-cells sorted into naive and memory

- T-cells sorted into naive CD4, naive CD8, memory CD4 and memory CD8
- Total of 60 repertoires: 20 B-cell repertoires and 40 T-cell repertoires

Walkthrough

We'll use the airr-standards docker image for this example, which comes loaded with all the python packages needed. You will want to map a local directory inside the docker container so you can access the data and analysis results afterwards:

```
# Download the image
docker pull airrc/airr-standards:latest
# Make local temporary directory to hold the data
mkdir adc_example
cd adc_example
# Invoke a shell session inside the docker image
docker run -it -v $PWD:/data airrc/airr-standards:latest bash
```

The first python script queries the data from the VDJServer data repository and saves them into files:

```
# Query the data
cd /data
python3 /airr-standards/docs/examples/api/retrieve_data.py
```

Only a subset of the data is downloaded for illustration purposes, but review the code to see how all data can be downloaded. A total of 40 repertoires and 300,178 rearrangements should be downloaded. The repertoire metadata is saved in the repertoires.airr.json file, and the rearrangements are saved in the rearrangements.tsv file. The script should take a few minutes to run and produce the following display messages:

```
Info: VDJServer Community Data Portal
    version: 1.3
description: VDJServer ADC API response for repertoire query
Received 40 repertoires.
Retrieving rearrangements for repertoire: 5168912186246295065-242ac11c-0001-012
Retrieved 9768 rearrangements for repertoire: 5168912186246295065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 5338391595746455065-242ac11c-0001-012
Retrieved 5521 rearrangements for repertoire: 5338391595746455065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 4858300151399575065-242ac11c-0001-012
Retrieved 2885 rearrangements for repertoire: 4858300151399575065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 5039977268020375065-242ac11c-0001-012
Retrieved 4053 rearrangements for repertoire: 5039977268020375065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 6240077029868695065-242ac11c-0001-012
Retrieved 3506 rearrangements for repertoire: 6240077029868695065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 6389112395039895065-242ac11c-0001-012
Retrieved 2289 rearrangements for repertoire: 6389112395039895065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 5939858815878295065-242ac11c-0001-012
Retrieved 3637 rearrangements for repertoire: 5939858815878295065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 6088937130722455065-242ac11c-0001-012
Retrieved 9028 rearrangements for repertoire: 6088937130722455065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 7446748091679895065-242ac11c-0001-012
Retrieved 1540 rearrangements for repertoire: 7446748091679895065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 7591789137265815065-242ac11c-0001-012
Retrieved 10000 rearrangements for repertoire: 7591789137265815065-242ac11c-0001-012
Retrieving rearrangements for repertoire: 7066128089908375065-242ac11c-0001-012
```

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Retrieved 5662 rearrangements for repertoire: 7066128089908375065-242ac11c-0001-012 Retrieving rearrangements for repertoire: 5624006920930455065-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 5624006920930455065-242ac11c-0001-012 Retrieving rearrangements for repertoire: 8961797805343895065-242ac11c-0001-012 Retrieved 1179 rearrangements for repertoire: 8961797805343895065-242ac11c-0001-012 Retrieving rearrangements for repertoire: 9084118473933975065-242ac11c-0001-012 Retrieved 4464 rearrangements for repertoire: 9084118473933975065-242ac11c-0001-012 Retrieving rearrangements for repertoire: 8485700680582295065-242ac11c-0001-012 Retrieved 3908 rearrangements for repertoire: 8485700680582295065-242ac11c-0001-012 Retrieving rearrangements for repertoire: 7309695685264535065-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 7309695685264535065-242ac11c-0001-012 Retrieving rearrangements for repertoire: 8425807333172056551-242ac11c-0001-012 Retrieved 6863 rearrangements for repertoire: 8425807333172056551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 8263242821018456551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 8263242821018456551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 8733756488295256551-242ac11c-0001-012 Retrieved 5298 rearrangements for repertoire: 8733756488295256551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 8602072790999896551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 8602072790999896551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 7313153105470296551-242ac11c-0001-012 Retrieved 9121 rearrangements for repertoire: 7313153105470296551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 6964444710708056551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 6964444710708056551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 7640859110155096551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 7640859110155096551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 7461458326201176551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 7461458326201176551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 5953881855632216551-242ac11c-0001-012 Retrieved 5916 rearrangements for repertoire: 5953881855632216551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 7158276584776536551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 7158276584776536551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 6393557657723736551-242ac11c-0001-012 Retrieved 7257 rearrangements for repertoire: 6393557657723736551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 6205695788196696551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 6205695788196696551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 4476756703191896551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 4476756703191896551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 4357957907784536551-242ac11c-0001-012 Retrieved 7033 rearrangements for repertoire: 4357957907784536551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 4931851437876056551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 4931851437876056551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 4744762662462296551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 4744762662462296551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 3252733973504856551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 3252733973504856551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 2989624276951896551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 2989624276951896551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 3628844259615576551-242ac11c-0001-012 Retrieved 5208 rearrangements for repertoire: 3628844259615576551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 3438706057421656551-242ac11c-0001-012 Retrieved 9530 rearrangements for repertoire: 3438706057421656551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 2197374609531736551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 2197374609531736551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 1993707260355416551-242ac11c-0001-012 Retrieved 10000 rearrangements for repertoire: 1993707260355416551-242ac11c-0001-012 Retrieving rearrangements for repertoire: 2541616238306136551-242ac11c-0001-012 Retrieved 6512 rearrangements for repertoire: 2541616238306136551-242ac11c-0001-012

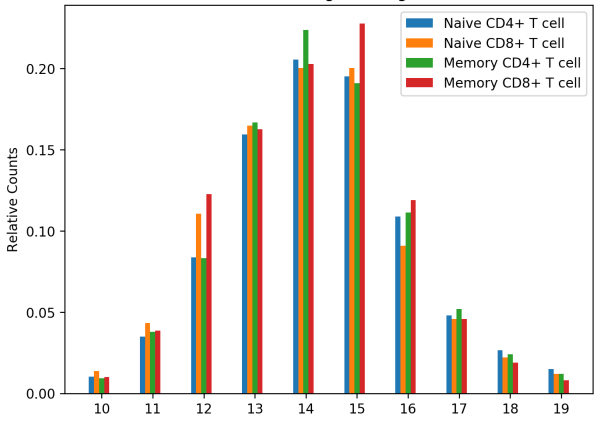
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```
Retrieving rearrangements for repertoire: 2366080924918616551-242ac11c-0001-012
Retrieved 10000 rearrangements for repertoire: 2366080924918616551-242ac11c-0001-012
```

The second python script loads the data from the files and generates a CDR3 amino acid length distribution that is grouped by the T cell subset. This study performs flow sorting to generate four T cell subsets: naive CD4+, naive CD8+, memory CD4+, memory CD8+. The script uses the repertoire metadata to determine the T cell subset for the rearrangement, tabulates the counts, normalizes them, and generates a grouped bar chart with the results:

```
# Run the analysis
python3 /airr-standards/docs/examples/api/analyze_data.py
```

The figure is placed in the plot.png file and should look like this:



CDR3 AA Length Histogram

Fig. 3: CDR3 AA Length Histogram grouped by T cell subsets.

Scientific Query Scenarios for AIRR Data Commons API

The AIRR Common Repository Working Group (CRWG) has defined a number of sample scientific query scenarios to guide the design of the ADC API. The Design Decisions document lists the major design choices for the API, and the *API* is currently defined using the OpenAPI V2.0 Specification. This document describes the query examples with associated JSON definitions that can be submitted to an AIRR repository.

There are two main query endpoints in the API: /repertoire for querying MiAIRR-compliant study metadata and /rearrangement for querying rearrangement annotations. Most scientific queries will involve both endpoints. The basic workflow involves first querying /repertoire to get the list of repertoires that meet the search criteria on study, subject, and sample metadata. Secondly, the identifiers from the repertoires in the first query are passed to the /rearrangement endpoint along with any search criteria on the rearrangement annotations. The resultant rearrangements can be downloaded as JSON or in the *AIRR TSV format*.

Query Example 1

What human full length TCR-beta sequences have junction amino acid sequence: "CASSYIKLN"?

• The JSON query definition for /repertoire endpoint. The ontology identifier 9606 requests human and TRB is the locus of interest.

```
{
    "filters":{
        "op":"and",
        "content": [
             {
                 "op":"=",
                 "content": {
                     "field":"subject.organism.id",
                     "value":"9606"
                 }
            },
             {
                 "op":"=",
                 "content": {
                     "field":"sample.pcr_target.pcr_target_locus",
                     "value":"TRB"
                 }
            }
        ]
    }
}
```

• That query does not request full length sequences. We can enhance the query by adding a clause for the sample.complete_sequences field.

```
{
    "filters":{
        "op":"and",
        "content": [{
            "op":"=",
            "content": {
                "field":"subject.organism.id",
                 "value":"9606"
            }
        },
        {
            "op":"=",
            "content": {
                "field":"sample.pcr_target.pcr_target_locus",
                 "value":"TRB"
            }
        },
        {
            "op":"or",
```

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```
"content":[{
                 "op":"=",
                 "content": {
                     "field":"sample.complete_sequences",
                     "value":"complete"
                 }
            },
            {
                 "op":"=",
                 "content": {
                     "field":"sample.complete_sequences",
                     "value": "complete+untemplated"
                 }
            }]
        }]
    }
}
```

• The JSON query definition for /rearrangement endpoint. The repertoire identifiers (repertoire_id) in the query are just examples, you would replace them with the actual identifiers returned from the above repertoire query. The query performs an exact match of the junction amino acid sequence.

```
{
    "filters":{
        "op":"and",
        "content": [
            {
                "op":"in",
                "content": {
                     "field": "repertoire_id",
                     "value":[
                         "2366080924918616551-242ac11c-0001-012",
                         "2541616238306136551-242ac11c-0001-012",
                         "1993707260355416551-242ac11c-0001-012",
                         "1841923116114776551-242ac11c-0001-012"
                     ]
                 }
            },
            {
                "op":"=",
                 "content": {
                     "field":"junction_aa",
                     "value": "CARDPRSYHAFDIW"
                 }
            }
        ]
    },
    "fields":["repertoire_id", "sequence_id", "v_call", "productive"],
    "format":"tsv"
```

Query Example 2

What human full length IgH sequences have been found in patients with an autoimmune diagnosis.

• TO BE WRITTEN

Query Example 3

What is the antibody IG heavy chain V usage in people who have diabetes?

• TO BE WRITTEN

Query Example 4

Give me all the anti-HIV antibody sequences that use IGHV1-69 in HIV infected individuals?

• TO BE WRITTEN

Query Example 5

Repertoires from cancer patients where we have pre- and post-immunotherapy peripheral blood (or tumor biopsy).

• TO BE WRITTEN

Query Example 6

Return TCRs that score highly on a position weight matrix from subjects with a particular HLA allele that have been infected with TB.

• TO BE WRITTEN

Query Example 7

Repertoires from female patients with cancer.

• TO BE WRITTEN

2.2 Release Notes

2.2.1 Schema Release Notes

Version 1.3.1: October 13, 2020

Version 1.3 documentation patch release.

Alignment Schema:

1. Added the deprecation tags for rearrangement_id, which were accidentally left out of the v1.3.0 release.

Version 1.3.0: May 28, 2020

Version 1.3 schema release.

New Schema:

- 1. Introduced the Repertoire Schema for describing study meta data.
- 2. Introduced the PCRTarget Schema for describing primer target locations.
- 3. Introduced the SampleProcessing Schema for describing experimental processing steps for a sample.
- 4. Replaced the SoftwareProcessing schema with the DataProcessing schema.
- 5. Introduced experimental schema for clonal clusters, lineage trees, tree nodes, and cells as Clone, Tree, Node, and Cell objects, respectively.

General Updates:

- 1. Added multiple additional attributes to a large number of schema propertes as AIRR extension attributes in the x-airr field. The new Attributes object contains definitions for these x-airr field attributes.
- 2. Added the top level required property to all relevant schema objects.
- 3. Added the title attribute containing the short, descriptive name to all relevant schema object fields.
- 4. Added an example attribute containing an example data value to multiple schema object fields.

AIRR Data Commons API:

1. Added OpenAPI V2 specification (specs/adc-api.yaml) for AIRR Data Commons API major version 1.

Ontology Support:

- 1. Added Ontology and CURIEResolution objects to support ontologies.
- 2. Added vocabularies/ontologies as JSON string for: Cell subset, Target substrate, Library generation method, Complete sequences, Physical linkage of different loci.

Rearrangement Schema:

- 1. Added the complete_vdj field to annotate whether a V(D)J alignment was full length.
- 2. Added the junction_length_aa field defining the length of the junction amino acid sequence.
- 3. Added the repertoire_id, sample_processing_id, and data_processing_id fields to serve as linkers to the appropriate metadata objects.
- 4. Added a controlled vocabulary to the locus field: IGH, IGI, IGK, IGL, TRA, TRB, TRD, TRG.
- 5. Deprecated the rearrangement_set_id and germline_database fields.
- 6. Deprecated rearrangement_id field and made the sequence_id field be the primary unique identifer for a rearrangement record, both in files and data repositories.
- 7. Added support secondary D gene rearrangement through the additional fields: d2_call, d2_score, d2_identity, d2_support, d2_cigar np3, np3_aa, np3_length, n3_length, p5d2_length, p3d2_length, d2_sequence_start, d2_sequence_end, d2_germline_start, d2_germline_start, d2_alignment_start, d2_alignment_end, d2_sequence_alignment, d2_sequence_alignment_aa, d2_germline_alignment_aa.
- 8. Updated field definitions with more concise V(D)J call descriptions.

Alignment Schema:

1. Deprecated the rearrangement_set_id and germline_database fields.

2. Added the data_processing_id field.

Study Schema:

1. Added the study_type field containing an ontology defined term for the study design.

Subject Schema:

- 1. Deprecated the organism field in favor of the new species field.
- 2. Deprecated the age field.
- 3. Introduced age ranges: age_min, age_max, and age_unit.

Diagnosis Schema:

1. Changed the type of the disease_diagnosis field from string to Ontology.

Sample Schema:

1. Changed the type of the tissue field from string to Ontology.

CellProcessing Schema:

- 1. Changed the type of the cell_subset field from string to Ontology.
- 2. Introduced the cell_species field which denotes the species from which the analyzed cells originate.

NucleicAcidProcessing Schema:

- 1. Defined the template_class field as type string.
- 2. Added a controlled vocabulary the library_generation_method field.
- 3. Changed the controlled vocabulary terms of complete_sequences. Replacing complete & untemplated with complete+untemplated and adding mixed.
- 4. Added the pcr_target field referencing the new PCRTarget schema object.

SequencingRun Schema:

- 1. Added the sequencing_run_id field which serves as the object identifer field.
- 2. Added the sequencing_files field which links to the RawSequenceData schema objects defining the raw read data.

RawSequenceData Schema:

- 1. Added the file_type field defining the sequence file type. This field is a controlled vocabulary restricted to: fasta, fastq.
- 2. Added the paired_read_length field defining mate-pair read lengths.
- 3. Defined the read_direction and paired_read_direction fields as type string.

DataProcessing Schema:

- 1. Replaces the SoftwareProcessing object.
- 2. Added data_processing_id, primary_annotation, data_processing_files, germline_database and analysis_provenance_id fields.

Version 1.2.1: Oct 5, 2018

Minor patch release.

1. Schema gene vs segment terminology corrections

- 2. Added Info object
- 3. Updated cell_subset URL in AIRR schema

Version 1.2.0: Aug 18, 2018

Peer reviewed released of the Rearrangement schema.

- 1. Definition change for the coordinate fields of the Rearrangement and Alignment schema. Coordinates are now defined as 1-based closed intervals, instead of 0-based half-open intervals (as previously defined in v1.1 of the schema).
- 2. Removed foreign study_id fields
- 3. Introduced keywords_study field

Version 1.1.0: May 3, 2018

Initial public released of the Rearrangement and Alignment schemas.

- 1. Added required and nullable constrains to AIRR schema.
- 2. Schema definitions for MiAIRR attributes and ontology.
- 3. Introduction of an x-airr object indicating if field is required by MiAIRR.
- 4. Rename rearrangement_set_id to data_processing_id.
- 5. Rename study_description to study_type.
- 6. Added physical_quantity format.
- 7. Raw sequencing files into separate schema object.
- 8. Rename Attributes object.
- 9. Added primary_annotation and repertoire_id.
- 10. Added diagnosis to repertoire object.
- 11. Added ontology for organism.
- 12. Added more detailed specification of sequencing_run, repertoire and rearrangement.
- 13. Added repertoire schema.
- 14. Rename definitions.yaml to airr-schema.yaml.
- 15. Removed c_call, c_score and c_cigar from required as this is not typical reference aligner output.
- 16. Renamed vdj_score, vdj_identity, vdj_evalue, and vdj_cigar to score, identity, evalue, and cigar.
- 17. Added missing c_identity and c_evalue fields to Rearrangement spec.
- 18. Swapped order of N and S operators in CIGAR string.
- 19. Some description clean up for consistency in Rearrangement spec.
- 20. Remove repeated objects in definitions.yaml.
- 21. Added Alignment object to definitions.yaml.
- 22. Updated MiARR format consistency check TSV with junction change.
- 23. Changed definition from functional to productive.

Version 1.0.1: Jan 9, 2018

MiAIRR v1 official release and initial draft of Rearrangement and Alignment schemas.

2.2.2 Python Library Release Notes

Version 1.3.1: October 13, 2020

- 1. Refactored merge_rearrangement to allow for larger number of files.
- 2. Improved error handling in format validation operations.

Version 1.3.0: May 30, 2020

- 1. Updated schema set to v1.3.
- 2. Added load_repertoire, write_repertoire, and validate_repertoire to airr. interface to read, write and validate Repertoire metadata, respectively.
- 3. Added repertoire_template to airr.interface which will return a complete repertoire object where all fields have null values.
- 4. Added validate_object to airr.schema that will validate a single repertoire object against the schema.
- 5. Extended the airr-tools commandline program to validate both rearrangement and repertoire files.

Version 1.2.1: October 5, 2018

1. Fixed a bug in the python reference library causing start coordinate values to be empty in some cases when writing data.

Version 1.2.0: August 17, 2018

- 1. Updated schema set to v1.2.
- 2. Several improvements to the validate_rearrangement function.
- 3. Changed behavior of all *airr.interface* functions to accept a file path (string) to a single Rearrangement TSV, instead of requiring a file handle as input.
- 4. Added base argument to RearrangementReader and RearrangementWriter to support optional conversion of 1-based closed intervals in the TSV to python-style 0-based half-open intervals. Defaults to conversion.
- 5. Added the custom exception ValidationError for handling validation checks.
- 6. Added the validate argument to RearrangementReader which will raise a ValidationError exception when reading files with missing required fields or invalid values for known field types.
- 7. Added validate argument to all type conversion methods in Schema, which will now raise a ValidationError exception for value that cannot be converted when set to True. When set False (default), the previous behavior of assigning None as the converted value is retained.
- Added validate_header and validate_row methods to Schema and removed validations methods from RearrangementReader.
- 9. Removed automatic closure of file handle upon reaching the iterator end in RearrangementReader.

Version 1.1.0: May 1, 2018

Initial release.

2.2.3 R Library Release Notes

Version 1.3.0: May 26, 2020

- 1. Updated schema set to v1.3.
- 2. Added info slot to Schema object containing general schema information.

Version 1.2.0: August 17, 2018

- 1. Updated schema set to v1.2.
- 2. Changed defaults to base="1" for read and write functions.
- 3. Updated example TSV file with coordinate changes, addition of germline_alignment data and simplification of sequence_id values.

Version 1.1.0: May 1, 2018

Initial release.

2.3 AIRR Standards

Information about all of the AIRR Community standards.

2.3.1 Introduction to MiAIRR

Summary

One of the primary initiatives of the Adaptive Immune Receptor Repertoire (AIRR) Community has been to develop a set of metadata standards for the submission of AIRR sequencing datasets. This work has been carried out by the AIRR Community Minimal Standards Working Group. In order to support reproducibility, standard quality control, and data deposition in a common repository, the AIRR Community has agreed to six high-level data sets that will guide the publication, curation and sharing of AIRR-Seq data and metadata: Study and subject, sample collection, sample processing and sequencing, raw sequences, processing of sequence data, and processed AIRR sequences. The detailed data elements within these sets are defined *here* (Download as TSV).

Topics

MiAIRR Data Elements

The AIRR Community has agreed to six high-level data sets that will guide the publication, curation and sharing of AIRR-Seq data and metadata: Study and subject, sample collection, sample processing and sequencing, raw sequences, processing of sequence data, and processed AIRR sequences.

Download as TSV.

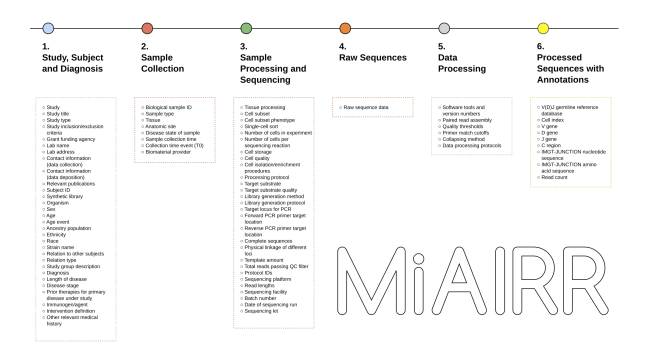


Fig. 4: Schema of MiAIRR data sets and the individual data elements of each set.

Set	Designation /	Type / For-	Level	Definition	Example
/	Field	mat			
Sub	-				
set					
1 /	Study ID	string free text	important	Unique ID assigned by study	PRJNA001
stud	ystudy_id			registry	
1 /	Study title	string free text	important	Descriptive study title	Effects of sun light
stud	ystudy_title				exposure of the Treg
					repertoire
1 /	Study type	Ontology	important	Type of study design	id: NCIT:C15197,
stud	ystudy_type	Ontology: {			value: Case-Control
		top_node:			Study
		{ id:			
		NCIT:C63536,			
		value:			
		Study}}			
1 /	Study inclu-	string free text	important	List of criteria for inclu-	Include: Clinical
stud	y sion/exclusion			sion/exclusion for the study	P. falciparum in-
	criteria				fection; Exclude:
	inclusion_e	xclusion_cr:	lteria		Seropositive for
					HIV
1 /	Grant fund-	string free text	important	Funding agencies and grant	NIH, award number
stud	y ing agency			numbers	R01GM987654
	grants				

	Desite a di di			nom previous page	
/	Designation / Field	Type / For- mat	Level	Definition	Example
Sub set	-				
1/	Contact infor-	string free text	important	Full contact information of the	Dr. P. Stibbons,
study	y mation (data			data collector, i.e. the person	p.stibbons@unseenu.edu
[collection)			who is legally responsible for	-
	collected_b	V		data collection and release. This	
	_	-		should include an e-mail ad-	
				dress.	
1/	Lab name	string free text	important	Department of data collector	Department for Pla-
stud	ylab_name	20	1	1	nar Immunology
1/	Lab address	string free text	important	Institution and institutional ad-	School of Medicine,
study	ylab_address		I	dress of data collector	Unseen University,
ĺ	, <u>-</u>				Ankh-Morpork,
					Disk World
1/	Contact infor-	string free text	important	Full contact information of the	Adrian Turnipseed,
	y mation (data		P or canto	data depositor, i.e. the person	a.turnipseed@unseenu.edu
Judy	deposition)			submitting the data to a repos-	
	submitted_b	177		itory. This is supposed to be	
	Submitteed_L	Y Y		a short-lived and technical role	
				until the submission is relased.	
1/	Relevant	string free text	important	Publications describing the ra-	PMID:85642
	y publications	sung free lexi	mportant	tionale and/or outcome of the	1 WILD:05042
study	pub_ids			study	
1/	Keywords	array of string	important	Keywords describing properties	['contains_ig',
study	•	<i>Controlled</i>	Important	of one or more data sets in a	'con-
study	keywords_st			study	tains_paired_chain']
	keywords_st	lary: con-		study	tams_paried_chain]
		tains_ig, con-			
		tains_tcr, con-			
		tains_icr, con- tains_single_cel	11		
			ι,		
		con-	ain.		
1/	Cubicat ID	tains_paired_ch		Subject ID agains of her sub with	SUD956412
1/	Subject ID	string free text	important	Subject ID assigned by submit-	SUB856413
sub-	subject_id			ter, unique within study	
ject	Sweth at -	hooloor tour	accontic1	TRUE for libraries in which the	
1/	Synthetic	boolean <i>true</i>	essential		
sub-	library	false		diversity has been synthetically	
ject	synthetic			generated (e.g. phage display)	· 1
1/	Organism	Ontology	essential	Binomial designation of sub-	id:
sub-	species	Ontology: {		ject's species	NCBITAXON:9606,
10.04					value: Homo sapi-
ject		top_node:			-
ject		{ id:			ens
ject		{ id: NCBITAXON:72	776,		-
jeci		{ id: NCBITAXON:72 value:	776,		-
ject		{ id: NCBITAXON:72	776,		-

Table 1 – continued from previous page

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Set	Designation /	Type / For-	Level	Definition	Example
/	Field	mat			
Sub	-				
set					
1 /	Sex sex	string Con-	important	Biological sex of subject	female
sub-		trolled vo-			
ject		cabulary:			
		male, fe-			
		male, pooled,			
		hermaphrodite,			
		intersex, not			
		collected, not			
		applicable			
1/	Age minimum	number posi-	important	Specific age or lower boundary	60
sub-	age_min	tive number		of age range.	
ject	ago				
1/	Age max-	number posi-	important	Upper boundary of age range	80
sub-	imum	tive number	mportunt	or equal to age_min for specific	
ject	age_max			age. This field should only be	
jeer	age_max			null if age_min is null.	
1 /	Age unit	Ontology	important	Unit of age range	id: UO:000036,
sub-	-		mportant	Unit of age range	· · ·
	age_unit	Ontology: {			value: year
ject		top_node:			
		{ id:			
		UO:000003,			
		value: time			
1 /	•	unit}}	• • •		11
1/	Age event	string free text	important	Event in the study schedule to	enrollment
sub-	age_event			which Age refers. For NCBI	
ject				BioSample this MUST be sam-	
				pling. For other implemen-	
				tations submitters need to be	
				aware that there is currently no	
				mechanism to encode to poten-	
				tial delta between Age event and	
				Sample collection time, hence	
				the chosen events should be in	
				temporal proximity.	
1 /	Ancestry	string free text	important	Broad geographic origin of an-	list of continents,
sub-	population			cestry (continent)	mixed or unknown
ject	ancestry_po	pulation			
1/	Ethnicity	string <i>free text</i>	important	Ethnic group of subject (de-	English, Kurds,
sub-	ethnicity		-	fined as cultural/language-based	Manchu, Yakuts
ject	-			membership)	(and other fields
				1/	from Wikipedia)
1/	Race race	string free text	important	Racial group of subject (as de-	White, American
sub-		0,	T	fined by NIH)	Indian or Alaska
ject					Native, Black,
,					Asian, Native
					Hawaiian or Other
					Pacific Islander,
					Other
					tinued on next nade

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Set	Designation /	Type / For-	Level	Definition	Example
/	Field	mat			
Sub					
set	-				
1/	Strain name	string free text	important	Non-human designation of the	C57BL/6J
sub-	strain_name			strain or breed of animal used	
ject	D 1 d		• • •		GUD 1055(40
1/	Relation to	string free text	important	Subject ID to which Relation	SUB1355648
sub-	other subjects			type refers	
ject	linked_subj		in a sut sut	Delation between subject and	fathan darahtan
1/	Relation type	string free text	important	Relation between subject and	father, daughter,
sub-	link_type			<i>linked_subjects</i> , can be genetic	household
ject	Study mayn	atring fue a tout	immontant	or environmental (e.g.exposure)	aantral
1 / di-	Study group description	string free text	important	Designation of study arm to which the subject is assigned to	control
	-	_description		which the subject is assigned to	
ag- no-	scuuy_group	_aescription	1		
sis					
and					
in-					
ter-					
ven-					
tion					
1/	Diagnosis	Ontology	important	Diagnosis of subject	id: DOID:9538,
di-	disease_dia	01	1		value: multiple
ag-		top_node: {			myeloma
no-		id: DOID:4,			·
sis		value: dis-			
and		ease}}			
in-					
ter-					
ven-					
tion					
1 /	Length of	string free text	important	Time duration between initial	23 months
di-	disease			diagnosis and current interven-	
ag-	disease_ler	lgth		tion	
no-					
sis					
and .					
in-					
ter-					
ven-					
tion				0.00	tinued on next page

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Set	Designation /	Type / For-	Level	Definition	Example
/	Field	mat			
Sub					
set					
1/	Disease stage	string free text	important	Stage of disease at current inter-	Stage II
di-	disease_sta		important	vention	Stugen
ag-	<u>arsease_</u> see	.90		(children	
no-					
sis					
and					
in-					
ter-					
ven-					
tion					
1 /	Prior thera-	string free text	important	List of all relevant previous ther-	melphalan/prednisone
di-	pies for pri-	sumg free ieni	mportant	apies applied to subject for treat-	merphatan/predmsone
ag-	mary disease			ment of <i>Diagnosis</i>	
no-	under study			ment of Diagnosis	
sis	prior_thera	nioc			
and	prior_cnere	ipies			
in-					
ter-					
ven-					
tion 1 /	Immunogen/age	ntring fue a tout	immortant	Antigen, vaccine or drug ap-	bortezomib
di-		insuring free text	important	plied to subject at this interven-	bonezonno
	immunogen			tion	
ag-				tion	
no- sis					
and					
in-					
ter-					
tion					
1 /	Intervention	string free text	important	Description of intervention	systemic
di-	definition	sumg free lext	mportant	Description of intervention	chemotherapy,
		n			
ag-	interventio	11			6 cycles, 1.25 mg/m2
no-					mg/mz
sis					
and					
in-					
ter-					
ven-					
tion					tinued on next page

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Set Designation /			Definition	Evampla
Set Designation / / Field	Type / For-	Level	Deminion	Example
,	mat			
Sub-				
set		• • •		
1 / Other rele-	string <i>free text</i>	important	Medical history of subject that is	MGUS, first diag-
di- vant med-			relevant to assess the course of	nosed 5 years prior
ag- ical history			disease and/or treatment	
no- medical_his	tory			
sis				
and				
in-				
ter-				
ven-				
tion				GLUD 50 44 5
2 / Biological	string <i>free text</i>	important	Sample ID assigned by submit-	SUP52415
sam-sample ID			ter, unique within study	
ple sample_id		•		D'
2 / Sample type	string free text	important	The way the sample was ob-	Biopsy
sam-sample_type	1		tained, e.g. fine-needle aspirate,	
ple			organ harvest, peripheral venous	
			puncture	
2 / Tissue	Ontology	important	The actual tissue sampled, e.g.	id:
sam-tissue	Ontology: {		lymph node, liver, peripheral	UBERON:0002371,
ple	top_node:		blood	value: bone marrow
	{ id:			
	UBERON:0010	<i>000</i> ,		
	value: mul-			
	ticellular			
	anatomical			
	structure}}			
2 / Anatomic site	string <i>free text</i>	important	The anatomic location of the tis-	Iliac crest
sam anatomic_si	te		sue, e.g. Inguinal, femur	
ple				
2 / Disease state	string free text	important	Histopathologic evaluation of	Tumor infiltration
sam of sample			the sample	
ple disease_sta	te_sample			
2 / Sample col-	string free text	important	Time point at which sample was	14 d
sam-lection time			taken, relative to Collection time	
ple collection_	time_point_:	relative	event	
2 / Collection	string free text	important	Event in the study schedule to	Primary vaccination
sam- time event			which Sample collection time	
ple collection_	time_point_:	reference	relates to	
2 / Biomaterial	string free text	important	Name and address of the entity	Tissues-R-Us,
sam- provider		· ·	providing the sample	Tampa, FL, USA
ple biomaterial	_provider			± ′ ′
3 / Tissue pro-	string free text	important	Enzymatic digestion and/or	Collagenase
pro- cessing		r	physical methods used to isolate	A/Dnase I di-
cess tissue_prod	essing		cells from sample	gested, followed by
(cell)			r	Percoll gradient
	1	1	1	- sicon Siduloni

			from previous page	_
Set Designation /	Type / For-	Level	Definition	Example
/ Field	mat			
Sub-				
set				
3 / Cell subset	Ontology	important	Commonly-used designation of	id: CL:0000972,
pro- cell_subset		I · · · ·	isolated cell population	value: class
cess	top_node:		isolated cell population	switched mem-
(cell)	{ id:			ory B cell
(cen)	, t			ory b cen
	CL:0000542,			
	value: lym-			
	phocyte}}			
3 / Cell sub-	string <i>free text</i>	important	List of cellular markers and their	CD19+ CD38+
pro- set phenotype			expression levels used to isolate	CD27+ IgM- IgD-
cess cell_phenot	уре		the cell population	
(cell)				
3 / Cell species	Ontology	defined	Binomial designation of the	id:
pro- cell_specie			species from which the ana-	NCBITAXON:9606,
cess	top_node:		lyzed cells originate. Typi-	value: Homo sapi-
(cell)	{ id:		cally, this value should be iden-	ens
(cen)		776		CIIS
	NCBITAXON:7	770,	tical to <i>species</i> , if which case	
	value:		it SHOULD NOT be set explic-	
	Gnathos-		itly. Howver, there are valid ex-	
	tomata}}		perimental setups in which the	
			two might differ, e.g. chimeric	
			animal models. If set, this key	
			will overwrite the species infor-	
			mation for all lower layers of the	
			schema.	
3 / Single-	boolean true	important	TRUE if single cells were iso-	
	false	Important	lated into separate compart-	
1	, v			
cess single_cell	4		ments	
(cell)				
3 / Number	integer posi-	important	Total number of cells that went	1000000
pro- of cells in	tive integer		into the experiment	
cess experiment				
(cell) cell_number	1			
3 / Number of	integer posi-	important	Number of cells for each biolog-	50000
pro- cells per	tive integer	_	ical replicate	
cess sequenc-				
(cell) ing reaction				
cells_per_r	feaction			
3 / Cell storage	boolean <i>true</i>	important	TRUE if cells were cryo-	True
U		mportant	preserved between isolation and	1100
pro- cell_storag	gaise		*	
cess			further processing	
(cell)				0000 1 1 111
3 / Cell quality	string free text	important	Relative amount of viable cells	90% viability as de-
pro- cell_qualit	У		after preparation and (if applica-	termined by 7-AAD
cess			ble) thawing	
(cell)				
I			0	tinued on port page

Table 1 – continued from previo	ous page
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Set Designation / / Field Sub- set	Type / For- mat	Level	Definition	Example
3 / Cell isolation pro- / enrichment cess procedure (cell) cell_isolat	string free text	important	Description of the procedure used for marker-based isolation or enrich cells	Cells were stained with fluorochrome labeled antibodies and then sorted on a FlowMerlin (CE) cytometer.
3 / Processing pro- protocol cess cell_proces (cell)	string free text	important	Description of the meth- ods applied to the sample including cell preparation/ iso- lation/enrichment and nucleic acid extraction. This should closely mirror the Materials and methods section in the manuscript.	Stimulated wih anti- CD3/anti-CD28
3 / Target pro- substrate cess template_cl (nu- cleic acid)	string Con- trolled vocab- addary: DNA, RNA	essential	The class of nucleic acid that was used as primary starting material for the following pro- cedures	RNA
3 / Target sub- pro- strate quality cess template_qu (nu- cleic acid)	string free text	important	Description and results of the quality control performed on the template material	RIN 9.2
3 / Template pro- amount cess template_an (nu- cleic acid)	string free text	important	Amount of template that went into the process	1000 ng

Table 1	- continued from	previous page
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Set	Designation /	Type / For-	Level	Definition	Example
	Field	mat	20101		Example
Sub		mat			
set	_				
	T '1	C.		<u>C</u>	DTT(. 1'
3 /	Library gener-	string Con-	essential	Generic type of library genera-	RT(oligo-
pro-	ation method	trolled vocab-		tion	dT)+TS(UMI)+PCR
cess	library_ger	le <i>ulat</i> yion <u>P</u> G R ,th			
(nu-		RT(RHP)+PCR			
cleic		RT(oligo-			
acid)	dT)+ PCR ,			
		RT(oligo-			
		dT)+ TS + PCR ,			
		RT(oligo-			
		dT)+ $TS(UMI)$ +	PCR,		
		RT(specific)+PC			
		RT(specific)+TS			
		RT(specific)+TS			
		RT(specific+UM)			
		RT(specific+UM)			
		RT(specific)+TS			
		other	,		
21	T :1		:	Description of another of	
3 /	Library gener-	string free text	important	Description of processes ap-	cDNA was gener-
pro-	ation protocol		_	plied to substrate to obtain a li-	ated using
cess	library_ger	eration_prot	tocol	brary that is ready for sequenc-	
(nu-				ing	
cleic					
acid)				
3 /	Protocol IDs	string free text	important	When using a library genera-	v2.1 (2016-09-15)
pro-	library_ger	eration_kit_	version	tion protocol from a commer-	
cess				cial provider, provide the proto-	
(nu-				col version number	
cleic	;				
acid					
	,				

Table 1-	continued from previou	s page
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Set	Designation /	Type / For-	Level	Definition	Example
/	Field	mat			
Sub-	-				
set					
3 /	Complete	string Con-	essential	To be considered <i>complete</i> , the	partial
pro-	sequences	trolled vo-		procedure used for library con-	
cess	complete_se	opanbalaneys:		struction MUST generate se-	
(nu-		partial, com-		quences that 1) include the first	
cleic		plete, com-		V gene codon that encodes the	
acid)		plete+untempla	ited,	mature polypeptide chain (i.e.	
		mixed		after the leader sequence) and 2)	
				include the last complete codon	
				of the J gene (i.e. 1 bp 5'	
				of the J->C splice site) and	
				3) provide sequence informa-	
				tion for all positions between 1)	
				and 2). To be considered <i>com</i> -	
				plete & untemplated, the sec-	
				tions of the sequences defined	
				in points 1) to 3) of the previ-	
				ous sentence MUST be untem-	
				plated, i.e. MUST NOT overlap	
				with the primers used in library	
				preparation. <i>mixed</i> should only	
				be used if the procedure used for	
				library construction will likely	
				produce multiple categories of	
				sequences in the given experi-	
				ment. It SHOULD NOT be used	
				as a replacement of a NULL	
				value.	tipued on poyt page

Table 1 – continued from previous page

		2		from previous page	
	Designation /	Type / For-	Level	Definition	Example
/	Field	mat			
Sub	-				
set					
3 /	Physical	string Con-	essential	In case an experimental setup	hetero_head-head
pro-	linkage of	trolled vocab-		is used that physically links nu-	
cess	different re-	ulary: none,		cleic acids derived from dis-	
(nu-	arrangements	hetero_head-		tinct Rearrangements before li-	
	physical_li	nhreade		brary preparation, this field de-	
acid)	hetero_tail-		scribes the mode of that link-	
		head, het-		age. All <i>hetero</i> _* terms indicate	
		ero_prelinked		that in case of paired-read se-	
		-		quencing, the two reads should	
				be expected to map to distinct	
				IG/TR loci. *_head-head refers	
				to techniques that link the 5'	
				ends of transcripts in a single-	
				cell context. *_ <i>tail-head</i> refers	
				to techniques that link the 3' end	
				of one transcript to the 5' end of	
				another one in a single-cell con-	
				text. This term does not provide	
				any information whether a con-	
				tinuous reading-frame between	
				the two is generated. *_pre-	
				<i>linked</i> refers to constructs in	
				which the linkage was already	
				present on the DNA level (e.g.	
				scFv).	
3 /	Target lo-	string Con-	important	Designation of the target locus.	IGK
	Target lo- cus for PCR	trolled vocab-	important	Note that this field uses a con-	IUK
pro-					
cess	pcr_target_			trolled vocubulary that is meant	
(nu-		IGI, IGK,		to provide a generic classifica-	
cleic	;	IGL, TRA,		tion of the locus, not necessarily	
acid	D.	TRB, TRD,		the correct designation accord-	
[pcr]		TRG		ing to a specific nomenclature.	
3 /	Forward PCR	string free text	important	Position of the most distal nu-	IGHV, +23
-	primer tar-			cleotide templated by the for-	
	get location			ward primer or primer mix	
(nu-		_primer_targ	get_location		
cleic					
acid					
[pcr]					
3 /	Reverse PCR	string free text	important	Position of the most proximal	IGHG, +57
pro-	-			nucleotide templated by the re-	
cess	get location			verse primer or primer mix	
(nu-	reverse_pcr	_primer_targ	et_location		
cleic	;				
acid					
[pcr])				
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Cat Designation /			Definition	Evennele
Set Designation /	Type / For-	Level	Definition	Example
/ Field	mat			
Sub-				
set				
3 / Batch number	string free text	important	ID of sequencing run assigned	160101_M01234
pro- sequencing		-	by the sequencing facility	
cess	1		- y yyy	
(se-				
quenc-				
ing)				
3 / Total reads	integer posi-	important	Number of usable reads for	10365118
pro- passing	tive integer		analysis	
cess QC filter				
		filter		
quenc-	<u></u>			
ing)		• • •		A1
3 / Sequencing	string free text	important	Designation of sequencing in-	Alumina LoSeq
pro- platform			strument used	1000
cess sequencing	platform			
(se-				
quenc-				
ing)				
3 / Sequencing	string free text	important	Name and address of sequenc-	Seqs-R-Us, Vancou-
	sumg free text	mportant	ing facility	ver, BC, Canada
· ·	Const 1 days		ing facility	ver, bC, Callada
cess sequencing	facility			
(se-				
quenc-				
ing)				
3 / Date of se-	string free text	important	Date of sequencing run	2016-12-16
pro- quencing run		-		
cess sequencing	run date			
(se-				
`				
quenc-				
ing)				
3 / Sequencing	string <i>free text</i>	important	Name, manufacturer, order and	FullSeq 600,
pro- kit			lot numbers of sequencing kit	Alumina,
cess sequencing	kit			#M123456C0,
(se-				789G1HK
quenc-				
ing)				
4.4	string Con-	important	File format for the raw reads or	
	-	important		
data quencing	trolled vocab-		sequences	
(raw data file type	ulary: fasta,			
reads)file_type	fastq			
4 / Raw se-	string free text	important	File name for the raw reads	MS10R-NMonson-
data quencing data			or sequences. The first file in	C7JR9_S1_R1_001.fastq
	1		paired-read sequencing.	
(raw file name			paned-read sequencing.	
(raw file name reads)filename			parted-read sequencing.	

Table 1	- continued from	previous page
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			lable	1 – continued	from previous page	
Set Desig	nation /		or- I	Level	Definition	Example
/ Field		mat				
Sub-						
set						
		U		important	Read direction for the raw rea	
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(raw		cabulary:			paired-read sequencing.	
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4 / Forwa		integer pa		important	Read length in bases for the fi	
pro- read	-	tive integer			file in paired-read sequencing	5
	_length					
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		string <i>free t</i>		important		C7JR9_S1_R2_001.fasto
data sequer (raw data fi					paired-read sequencing	C/JK9_31_K2_001.1astq
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quenc-						
ing)						
5 / Softwa	are	string free t	ext i	important	Version number and / or da	ate, IgBLAST 1.6
pro- tools a	and ver-	0.5		1	include company pipelines	
cess sion	numbers					
(com-soft	ware_ve	rsions				
pu-						
ta-						
tional)						
5 / Paired	read	string free t	<i>ext</i> i	important	How paired end reads were	÷ .
pro- assem	bly				sembled into a single recep	-
cess pair	ed_reads	s_assembl	ly		sequence	old 0.8)
(com-						
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tional)						
5 / Qualit		string free t	<i>ext</i> i	important	How sequences were remov	-
pro- thresh					from (4) based on base qual	lity >=20
	ity_thre	esholds			scores	
(com-						
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I to I						
ta- tional)						

Table 1 – continued from previous page

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Set	Designation /	Type / For-	Level	Definition	Example
/	Field	mat			
Sub	-				
set					
5 /	Primer match	string free text	important	How primers were identified in	Hamming distance
pro-	cutoffs			the sequences, were they re-	<= 2
cess	primer_mato	h_cutoffs		moved/masked/etc?	
(con	1-				
pu-					
ta-					
tion	al)				
5 /	Collapsing	string free text	important	The method used for combining	MUSCLE 3.8.31
pro-	method			multiple sequences from (4) into	
cess	collapsing_	method		a single sequence in (5)	
(con	1-				
pu-					
ta-					
tion	/				
5 /	Data process-	string <i>free text</i>	important	General description of how QC	Data was processed
pro-	0 1			is performed	using []
cess	data_proces	sing_protoco	pls		
(con	1-				
pu-					
ta-					
tiona	,				
5 /	V(D)J	string free text	important	Source of germline V(D)J genes	ENSEMBL, Homo
	germline			with version number or date ac-	sapiens build 90,
· •	- reference			cessed.	2017-10-01
cess	edlatabase				
se-	germline_da	tabase			
quer	nce)				

Table 1 – continued from previous page

MiAIRR-to-NCBI Implementation

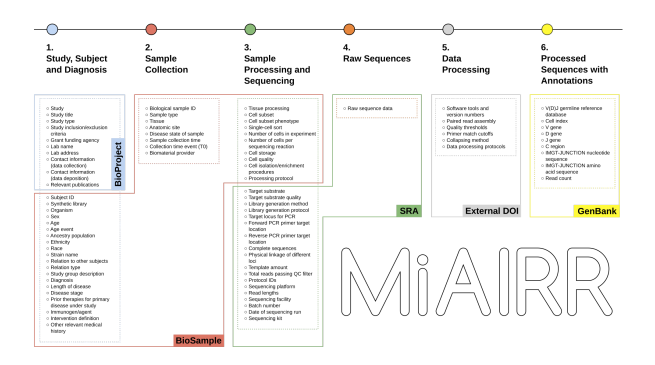
Authors Christian E. Busse, Florian Rubelt and Syed Ahmad Chan Bukhari

Guide for submission of AIRR-seq data to NCBI

This site provides a detailed "how-to" guide for submission of AIRR-seq data to **NCBI repositories** (BioProject, BioSample, SRA and GenBank). For other implementations of the MiAIRR standard see here.

One of the primary initiatives of the AIRR (Adaptive Immune Receptor Repertoire) Community has been to develop a set of metadata standards for the submission of immune receptor repertoire sequencing datasets. This work has been carried out by the AIRR Community Standards Working Group. In order to support reproducibility, standard quality control, and data deposition in a common repository, the AIRR Community has agreed to six high-level data sets that will guide the publication, curation and sharing of AIRR-Seq data and metadata: Study and subject, sample collection, sample processing and sequencing, raw sequences, processing of sequence data, and processed AIRR sequences. The detailed data elements within these sets are defined *here* (Download as TSV). The association between these AIRR sets, the associated data elements, and each of the NCBI repositories is shown below:

Submission of AIRR sequencing data and metadata to NCBI's public data repositories consists of five sequential steps:



- 1. Submit study information to NCBI BioProject using the NCBI web interface.
- 2. Submit sample-level information to the NCBI BioSample repository using the AIRR-BioSample templates.
- 3. Submit raw sequencing data to NCBI SRA using the AIRR-SRA data templates.
- 4. Generate a DOI for the protocol describing how raw sequencing data were processed using Zenodo.
- 5. Submit processed sequencing data with sequence-level annotations to GenBank using AIRR feature tags.

The submission manual provides step-by-step instructions on carrying out these steps for an AIRR study submission.

MiAIRR-to-NCBI Submission Manual

Scope of this document

Provide a user manual describing the submission of AIRR data using the NCBI reference implementation described in [Rubelt_2017]. This implementation uses NCBI's BioProject, BioSample, Sequence Read Archive (SRA) and GenBank repositories and metadata standards to report AIRR data.

Step 1. MiAIRR data submission to BioProject, BioSample and SRA

Since we propose to include a combination of raw and processed sequence data, the AIRR standard will sometimes need to be distributed and linked across multiple repositories (e.g., data in SRA linked to related data in GenBank). Besides, the data elements that comprise the standard will be mapped to ontologies in BioPortal through NIH CDE (Common Data Element) terms. These linkages will support more sophisticated validation and logical inference.

There are three main alternatives to submit raw AIRR data/metadata to NCBI repositories: (1) CEDAR's CAIRR pipeline; (2) NCBI's Web interface; and (3) NCBI's FTP server. These alternatives are described below:

Option 1. Submission via the CEDAR system (CAIRR submission pipeline)

CEDAR's CAIRR submission pipeline helps investigators and curators to edit and validate ontology-controlled metadata. This pipeline provides a seamless interface to transmit SRA datasets to the NCBI SRA and BioSample repositories from the CEDAR Workbench. The pipeline can be directly be accessed at http://cairr.airr-community.org. Note that the CEDAR template and template elements used by this pipeline are publicly available in the following CEDAR folder: All/Shared/Shared by CEDAR/MiAIRR.

Submission steps:

1. Open CEDAR's MiAIRR template by clicking on http://cairr.airr-community.org. If you are not already logged in, this will take you to the CEDAR login panel. If you are a new user, you will have to create an account on the CEDAR Workbench by clicking here.

LEDAR	
Email	
Password	
Remember me Forgot Password?	
LOG IN	
New user? Register g Google+ Image: Comparison of the second se	
Watch the video tutorial	

2. After logging in into the system, you will see the 'MiAIRR' template. Fill out the template fields with your metadata. Fields with an asterisk (*) are mandatory. Your submission will fail if any mandatory fields are not completed. If information is unavailable for any mandatory field, please enter 'not collected', 'not applicable' or 'missing' as appropriate. Note that you will need to enter a BioProject ID into the field 'Study ID'. If you do not have a BioProject yet, you can create one at *https://submit.ncbi.nlm.nih.gov/subs/bioproject/*

← 🔊 MiAIRR	• • •
MiAIRR	
Submissions Release Date	
BioProject for AIRR NCBI	
Study ID*	
Study Title*	
Study Type*	
Study Criteria	
Funding Agency*	
Contact Information (data collection)*	

3. Once your metadata is complete, click on the 'Save' button to save your metadata into your workspace. You will see a message in a green box confirming that your metadata have been successfully saved, as well as a message in a yellow box letting you know that your metadata have been saved to your personal workspace.

🗲 💽 MiAIR	R metadata			•••
	Batch Number*			
	AP626			
	Date of Sequencing Run*			
	5/3/2016			
	Sequencing Kit*			
	New England Biolabs			
	File Type			
	sra-run-fastq			
	filename			
	pass_nophix.fastq,	_001_ATCACG_L001_R1 _001_ATCACG_L001_R2		
	CANCEL	VALIDATE	SAVE	

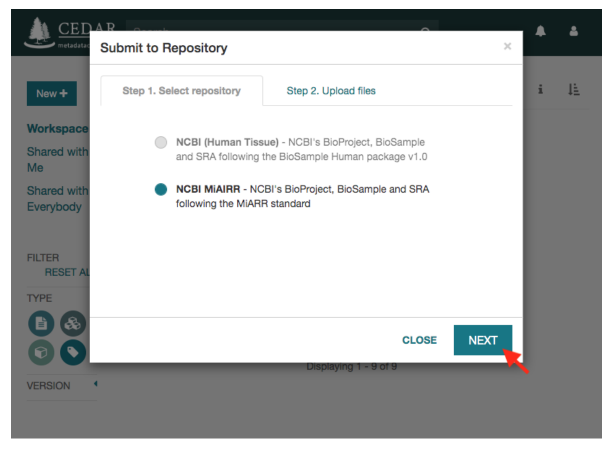
- 4. Go to your personal workspace by clicking on the left arrow (top left corner) and then on the 'Workspace' link, or by just clicking on: https://cedar.metadatacenter.org
- 5. Once in your workspace, you will see a metadata file called 'MiAIRR metadata'. That file contains the metadata that you have just created and that you want to submit to the NCBI. Click on the three vertical dots on the top-right corner of the file icon to see the available file options.

<u>CEDAR</u> metadatacenter.org	Search		Q		•	•
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Workspace Shared with Me Shared with Everybody	Field search demo	BioSample	PRECLINICALTRIAL			
FILTER RESET ALL TYPE TYPE	NCBI BioSample - H	Copy of NCBI BioSa Displaying 1	BioCADDIE metadata			

6. Click on the 'Submit' option to open the submission dialog.

				۹		•	•
New +	All / Users / Demo User				≔	i	μE
Workspace Shared with Me	Field search demo	BioSample					
Shared with Everybody	MiAIRR metadata	Populate Open	ia	PRECLINICALTRIAL			
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VERSION		Publish version Create version Submit	ring 1	- 9 of 9			
		Delete					

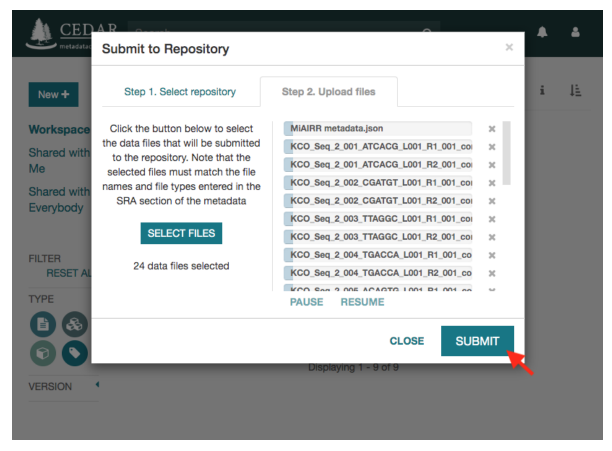
7. The 'NCBI MiAIRR' option will be automatically selected. Click on 'Next' to go to the next step.



8. Click on the 'Select Files' button to upload the data files. Note that the names of the selected files must match the names in the metadata file. Otherwise, you will receive an error message when trying to start the submission.

<u>CED</u> metadatac	A R Lourner Submit to Repository		0	×	•	۵
New +	Step 1. Select repository	Step 2. Upload files			i	1E
Workspace Shared with Me Shared with Everybody	Click the button below to select the data files that will be submitted to the repository. Note that the selected files must match the file names and file types entered in the SRA section of the metadata	MiAIRR metadata.json	1	×		
ТҮРЕ		PAUSE RESUME				
		Displaying 1 - 9	CLOSE	SUBMIT		
VERSION						

9. Click on the 'Submit' button to start the submission. If there are not validation errors, the selected data files and the corresponding metadata will be uploaded to the NCBI servers.



10. Note that the submission may take several hours or even days to be processed by the NCBI. Meanwhile, you will receive status messages about your submission in your workspace (messages icon).

<u>CEDAR</u> metadatacenter.or			Q		Messages	4
New +	All / Users / Demo User			≔	i	μ
Workspace Shared with Me	Field search demo	BioSample				
Shared with Everybody	MiAIRR metadata	CEDAR-NCBI Huma	PRECLINICALTRIAL			
FILTER RESET ALL	NCBI BioSample - H	Copy of NCBI BioSa	BioCADDIE metadata			
VERSION		Displaying 1	- 9 of 9			

11. Proceed with deposit of processed data, below.

Citing the CAIRR pipeline

Bukhari, Syed Ahmad Chan, Martin J. O'Connor, Marcos Martínez-Romero, Attila L. Egyedi, Debra Debra Willrett, John Graybeal, Mark A. Musen, Florian Rubelt, Kei H. Cheung, and Steven H. Kleinstein. The CAIRR pipeline for submitting standards-compliant B and T cell receptor repertoire sequencing studies to the NCBI. Frontiers in Immunology 9 (2018): 1877. DOI: 10.3389/fimmu.2018.01877

Tell Us About It

Please let us know how it went! If you are willing, we would love to have your comments in a short survey, it should just take 5 minutes or so. We also welcome entry of issues and requests in our GitHub repository, and emails can be sent to cedar-users@lists.stanford.edu. Both of these resources are publicly visible.

Support or Contact

Having trouble with NCBI submission process through our pipeline? Please email to Syed Ahmad Chan Bukhari or to Marcos Martínez-Romero and we will help you sort it out.

Option 2. Submission via NCBI's web interface

To facilitate AIRR data submissions to NCBI repositories, we have developed the NCBI-compliant metadata submission templates both for single and bulk AIRR data submissions. NCBI provides a web-based interface to create a BioProject and allows to BioSample, Sequence Read Archive (SRA) and GenBank metadata via tab-delimited files for single BioProject related data files submission.

Submitting AIRR data and associated metadata to the Bioproject, BioSample and SRA repositories via NCBI's web interface follows in general the submission procedure described in [NCBI_NBK47528], but uses AIRR-specific template for metadata submission:

- 1. Go to https://submit.ncbi.nlm.nih.gov/subs/sra/ and login with your NCBI account (create an account if necessary).
- 2. Click on "create new submission". You will see a form as below. Fill the form with required information and click on "continue".

NIH U.S. National Library of Medic	ine NCBI National Center for Biotechnology Information	ahmadchan@gmail.com
Submission Porta	номе	MY SUBMISSIONS GROUPS TEM
Sequence Read Archi test project, Dec 05 '16	ve (SRA) submission: SUB2154468	
1 SUBMITTER 2 GENERAL INF	• 3 project info 4 publications 5 biosample type 6 biosample attributes 7 sra metadata 8 files 9 o	VERVIEW
Submitter		Required fields are r
* First (given) name Middle nam	E-mail (secondary) • At least one e-mail should be from the organization's domain.	
Select group for this submission O None (affiliation from my person		
O Center for Expanded Data Annot	ation and Retrieval	
* Submitting organization	Submitting organization URL * Department	
Phone Fax P		
* Street	City State/Province Postal code * Country United States of America	

Continue SUpdate my contact information in profile

3. If you are submitting for the first time, check "Yes" on the "new BioProject" or "new BioSample" options to create a new project or sample, respectively.

Submission Portal		HOME	MY SUBMISSIONS	GROUPS	TEMPLATES	MY PRO
Sequence Read Archive (SRA) submission: SUB2154468 test project, Dec 05 '16					Delete	submis
1 SUBMITTER 2 CENERAL INFO 3 PROJECT INFO 4 PUBLICATIONS 5 BIOSAMPLE TYPE	26 biosample attributes 27 sra metadata 28	FILES 9 O	VERVIEW			
General Information			🙁 Rec	quired fields	are marked wi	ith aster
Do yog want to create new BioProject? ⊛ Yes ◎ No						
Do yog want to create new BioSamples for this submission? ⊛ Yes ○ No						
Release date						
 When should this submission be released to the public: Release immediately following processing (recommended) Release on specified date or upon publication, whichever is first Note: Release of BioProject or BioSample is also triggered by the release of linked data. 						
Continue						

4. Fill in the project information. Add as much relevant information you can add in description. It will help later in searching the particular submission.

1 JOBATTER 2 GENERAL INTO		3 3 MA MELADATA _ 0 THES _ 1 OT	2 C C C C C C C C C C C C C C C C C C C
Project Info			● R
* Project title 🔾			
* Public description 💿			
Relevance •		Fill project info	
* Is your project part of a larger init • No O Yes (not very common)	tiative which is already registered w	ith NCBI?	
External Links			
Link description 📀	URL 📀	Delete	
Add another link		•	
Select your grants			
 Use this tool to look up grants from NIH, CDC, FDA and VA) and some no Speaks). You can search by grant nu 	on-governmental funding sources (eg		
O Add grants			
Consortium name 🥥	Consortium URL 💿		
Data provider 오	Data provider URL 😡	Delete	

- 5. The AIRR BioSample template is not yet listed on the NCBI website. The template sheet AIRR_BioSample_V1.0.xls can be downloaded from https://github.com/airr-community/airr-standards/ tree/master/NCBI_implementation/templates_XLS. Fill in the required field and save the file as *tab-delimited* text file (.TSV format), then upload it.
- 6. To submit the SRA metadata use the AIRR_SRA_v1.0.xls file. Make sure that the column sample_name uses sample names that match the record in the BioSample template (if new BioSamples are being submitted) or a previously entered record. Also this file must be saved as *tab-delimited* text file for upload.
- 7. Submit the raw sequence file.
- 8. Complete the submission.
- 9. Proceed with deposit of processed data, below.

Option 3. Submission via NCBI's FTP server, using a predefined XML template

In addition to the web interface, NCBI provides an FTP-based solution to submit bulk metadata. The corresponding AIRR XML templates can be found under https://github.com/airr-community/airr-standards/tree/master/ NCBI_implementation/templates_XLS. Otherwise users should refer to the current SRA file upload manual https: //www.ncbi.nlm.nih.gov/sra/docs/submitfiles/. Users planning to frequently submit AIRR-seq data to SRA using scripts to generate the XML files MUST ensure that the templates are identical to the current upstream version on Github.

Step 2. Processed MiAIRR data submission to GenBank/TLS

Processed sequence data will be submitted to the "Targeted Locus Study" (TLS) section of GenBank. The details of this submission process are currently still being finalized. Basically the procedure is identical to a conventional GenBank submission with the exception of additional keywords marking it as TLS submission.

Non-productive records should be removed before the data submission or use an alternative annotation as described in the specification document.

• Generating MiAIRR compliant GenBank/TLS submissions: https://changeo.readthedocs.io/en/stable/examples/genbank.html

GenBank provides multiple tools (GUI and command-line) to submit data:

- BankIt, a web-based submission tool with wizards to guide the submission process
- Sequin, NCBI's stand-alone submission tool with wizards to guide the submission process is available by FTP for use on for Windows, macOS and Unix platforms.
- Tbl2asn is the recommended tool for the bulk data submission. It is a command-line program that automates the creation of sequence records files (.sqn) for submission to GenBank, driven by multiple tabular unput data files. Documentation and download options can be found under https://www.ncbi.nlm.nih.gov/genbank/tbl2asn2/.

MiAIRR-to-NCBI Specification

Outline of INSDC reporting procedure

TODO: Outline the reporting procedure for data sets 1-4

In terms of standard compliance it is currently REQUIRED¹ to deposit information for MiAIRR data sets 5 and 6 in general-purpose sequence repositories for which an AIRR-accepted specification on information mapping MUST exist. However, users should note that in the future additional AIRR-sanctioned mechanisms for data deposition will become available as specified by the AIRR Common Repository Working Group. The mapping of data items in MiAIRR data sets 5 and 6 differs substantially in size and structure and therefore requires distinct reporting procedures:

- Set 5: This is free text information describing the work flow, tools and parameters of the sequence read processing. It is REQUIRED that this information is deposited as a freely available document, permanently linked via a DOI. Note that is currently neither a specific format for this document nor a recommended service provider for obtaining the DOI.
- Set 6: This is specified to contain the consensus sequence and the following information obtained from the initial analysis: V, D and J segment, C region and IMGT-JUNCTION² [LIGMDB_V12]. These will be deposited in a general-purpose INSDC repository, using the record structure described below.

INSDC records were originally designed to hold individual Sanger sequences. Therefore each record will contain a header with information largely identical between all records in an AIRR sequencing study. Records can be concatenated for uploading.

The INSDC feature table (FT) [INSDC_FT] is a sequence annotation standard used within the INSDC records and assigns information to specified positions on the reported sequence string. In regard to the correct location of the provided annotation, it should especially be noted that some V(D)J inference tools will return coordinates referring to the reference instead of the query sequence. As the sequence submitted in a record MUST be identical to the query sequence, the positions provided by the V(D)J inference tool MUST, if necessary, be translated back onto the query sequence. In case the start and/or end of a feature cannot be reliably determined or is not present in the reported sequence³, open intervals CAN be used for reporting. However, open intervals MUST NOT be used to deliberately obfuscate known positions.

In addition to the required information specified in *Table_1*, users CAN use all valid FT keys/qualifiers to provide further annotation for the reported sequences. However, a record MUST still be compliant with this specification, if such OPTIONAL information would be removed, meaning that it is FORBIDDEN to move REQUIRED information into OPTIONAL keys/qualifiers. In addition, users MUST NOT use keys/qualifiers that could create ambiguity with the keys/qualifiers specified here.

¹ See the "Glossary" section on how to interpret term written in all-caps.

² Note that according to IMGT definition this is a superset of the CDR3.

³ This can occur e.g. in paired-end sequencing of head-to-head concatenated transcripts, where the 5' end of the V segment is present in the amplicon, but cannot be precisely determined.

element	FT key	FT qualifier	FT value	REQUIRED (if used by original study)
V segment	V_segment	/gene	see [Feature table]	yes
D segment	D_segment	/gene	see [Feature table]	yes; if IGH, TRB or TRD sequence
J segment	J_segment	/gene	see [Feature table]	yes
C region	C_region	/gene	see [Feature table]	yes
JUNCTION	CDS	/function	"JUNCTION"	yes

Table 1: Summary of the mapping of mandatory AIRR MiniStd data set 6 elements to the INSDC feature table (FT). Note that the overall record will contain additional information, such as cross-references linking the deposited sequence reads and metadata.

Element mapping

The broad strategy of element mapping to the various repositories is depicted in *Table_2*.

MiAIRR data set / subset	target repository
1 / study	BioProject
1 / subject	
1 / diagnosis & treatment	
2 / sample	BioSample
3 / processing (cells)	
3 / processing (nucleic acids)	SRA
4 / raw sequences	
5 / processing (data)	user-defined DOI
6 / Processed sequences & annotations	Genbank

Table 2: Summary of the mapping of MiAIRR data sets to the various repositories

Mapping of data sets 1-4 to BioProject/BioSample/SRA

TODO: Include item-by-item mapping [NCBI_NBK47528]

Mapping of data set 5 to a user-defined repository

While several mandatory item have been defined in this data set, there is currently no mapping as the reporting procedure is implemented as a free text document. AIRR RECOMMENDS to use Zenodo for deposition of these documents, as it is hosted by CERN and supports versioned DOIs (termed "concept" DOI). Users SHOULD use the existing AIRR tag when submitting documents to increase the visibility of their study.

Mapping of data set 6 to INSDC

Users should note that while the FT is standardized, the overall sequence record structure diverges between the three INSDC repositories. The following section refers to items at or above the hierarchy level of the FT using the GenBank specification [GENBANK_FF], the corresponding designations of ENA [ENA_MANUAL] are provided in parenthe-sis¹¹.

¹¹ Note that there is currently no submission specification for ENA. This information is provided for reference only and will be moved to a separate document in the future.

Record header

The header MUST contain all of the following elements:

- REQUIRED: header structure as specified by the respective INSDC repository [ENA_MANUAL] [GENBANK_FF] [GENBANK_SR].
- FORBIDDEN: The DEFINITION entry will be autopopulated by information provided in the FT part (misc_feature, /note).
- REQUIRED: identifier of the associated SRA record (MiAIRR data set 4) as DBLINK (ENA: DR line). Note that it is **not** possible to refer to individual raw reads, only the full SRA collections can be linked.
- REQUIRED: in the KEYWORDS field (ENA: KW line):
 - the term "TLS"
 - the term "Targeted Locus Study"
 - the term "AIRR"
 - the term "MiAIRR:<x>.<y>" with <x> and <y> indicating the used version and subversion of the MiAIRR standard.
- REQUIRED: DOI of the associated free-text record containing the information on data processing (MiAIRR data set 5) as REMARK within a REFERENCE⁴ (ENA: RX line).
- OPTIONAL: The use of structured comments is currently evalutated for use in future versions of the MiAIRR standard.

Feature table

The feature table, indicated by FEATURES (ENA: RX line), MUST or SHOULD contain the following keys/qualifiers:

General sequence information

- REQUIRED: key source containing the following qualifiers:
 - REQUIRED: qualifier /organism (required by [INSDC_FT]).
 - REQUIRED: qualifier /mol_type (required by [INSDC_FT]).
 - REQUIRED: qualifier /citation pointing to the reference in the header (REFERENCE, ENA: RN line) that links to the data set 5 document.
 - **REQUIRED**: qualifier / rearranged⁵.
 - REQUIRED: qualifier /note containing the AIRR_READ_COUNT keyword to indicate the read number used for the consensus. The criteria for selecting these reads and the procedure used to build the consensus SHOULD be reported as part of data set 5.
 - OPTIONAL: qualifier /note containing the AIRR_INDEX_CELL keyword for single-cell experiments. The value of the keyword SHOULD only contain alpha-numeric characters and MUST be identical for sequences derived from the same cell of origin.

⁴ The current GenBank record specification does not include a separate key for DOIs.

⁵ Although FT does specify a */germline* qualifier for non-rearranged sequences it has not been included in this specification as there is no obvious use case for it. In addition, non-rearranged transcripts would lack a number of other features that are assumed to be present, first of all the JUNCTION.

- RECOMMENDED: qualifiers /assembly_gap and /linkage_evidence to annotate nonoverlapping paired-end sequences.
- RECOMMENDED: qualifier / strain, if /organism is "Mus musculus".

Note that additional qualifiers might be REQUIRED by GenBank to harmonize the GenBank record with the BioSample referenced by it in the header. A list of known BioSample keyword and GenBank qualifiers that MUST contain the same information can be found below. Whether (and in which direction) the existence of a keyword/qualifiers triggers a requirement in the corresponding record is currently unknown. Please report any undocumented requirements surfacing during submission to the MiAIRR team.

BioSample keyword	GenBank FT qualifier
cell type	/cell_type
isolate	/isolate
sex	/sex
tissue	/tissue_type

Segment and region annotation

The following keys MUST be used for annotation according to their FT definition, if the respective item has been reported by the original study:

- REQUIRED: key V_region. Note that this key MUST NOT be used to annotate V segment leader sequence⁶⁷.
- REQUIRED: key misc_feature with coordinates identical to those given in V_region. This key MUST contain a /note qualifier that contains a string as value, which describes the general type of variable region described by the record. The string MUST match the regular expression

This string will be used as record heading upon import into Genbank. Note that while this behavior of Genbank is undocumented, the procedure has been approved by NCBI.

- REQUIRED: key V_segment, both coordinates MUST be within V_region. Note that this key MUST NOT be used to annotate V segment leader sequence⁶⁷.
- REQUIRED: key D_segment, both coordinates MUST be within V_region. This key is only REQUIRED for sequences of applicable loci (*IGH*, *TRB*, *TRD*⁸). In the rare case of rearrangements using two D segments, this key SHOULD occur twice, but the coordinates of both keys MUST NOT overlap.
- REQUIRED: key J_segment, both coordinates MUST be within V_region.
- REQUIRED: key C_region, both coordinates MUST NOT overlap with V_region. If the region can be unambiguously identified, the respective official gene symbol MUST be reported using the /gene qualifier. If only the isotype (e.g. IgG) but not the subclass (e.g. IgG1) can be identified, a truncated gene symbol (e.g. IGHG instead of IGHG1) SHOULD be reported instead⁹.

Each [VDJ]_segment key MUST or SHOULD contain the following qualifiers:

⁶ The FT explicitly states that $V_{segment}$ does **not** cover the leader sequence. The definition of V_{region} is slightly more ambiguous, however in combination with the $V_{segment}$ definition, it becomes clear that the leader is also not considered to be a part of V_{region} . Therefore the leader sequence should be implicitly annotated as the region between the start of *CDS* and the start of V_{region} .

⁷ Previously the leader was implicitly annotated as the region between *CDS* start and *V_region* start. As it was decided to drop the "global" CDS to make it easier to accommodate for INDELs, this is currently not an option anymore.

⁸ For simplicity, this document only uses human gene symbols. For non-human species the specification pertains to the respective orthologs.

⁹ This approach has been approved by NCBI.

- REQUIRED: qualifier /gene, containing the designation of the inferred segment, according to the database in the first /db_xref entry. This qualifier MUST NOT contain any allele information.
- RECOMMENDED: qualifier /allele, containing the designation of the inferred allele, according to the database in the first /db_xref entry. Note that while INSDC does not specify any format for this qualifier, AIRR compliance REQUIRES that this field only contains the allele string, i.e. without the gene name or separator characters.
- REQUIRED: qualifier /db_xref, linking to the reference record of the inferred segment in a germline database [INSDC_XREF]. This qualifier can be present multiple times, however only the first entry is mandatory and MUST link to the database used for the segment designation given with /gene and (if present) /allele.

Note on referencing IMGT databases: There are two IMGT database available in the controlled vocabulary [INSDC_XREF]:

- IMGT/GENE-DB: This is the genome database, which requires that a reference sequence has been mapped to genomic DNA. When using this database as reference, note that you can only refer to the gene symbol **not** the allele. In the case of ambiguous allele calls (see below) this means that you MUST NOT annotate any /allele at all. Nevertheless, this SHOULD be the default database for applications using IMGT as reference, as the sequence for each gene/allele is unique.
- IMGT/LIGM: This database collects sequences described in INSDC databases (GenBank/ENA/DDBJ). As it might contain multiple entries representing a given gene/allele, it is NOT RECOMMENDED to use it unless that inference gene/allele is only present in IMGT/LIGM and not in IMGT/GENE-DB.
- RECOMMENDED: /inference to indicate the tool used for segment inference. The description string SHOULD use COORDINATES as category and aligment as type [INSDC_FT].

Annotation of sequences producing multiple hits with identical scores is problematic and is ultimately at the discretion of the depositing researcher. However, the algorithms used for tie-breaking SHOULD be documented in data set 5. In addition, the following procedures MUST be followed:

- Certain gene, ambiguous allele: If multiple alleles of the same gene match to the sequence, the /allele qualifier MUST NOT be used. As the REQUIRED /db_xref qualifier will ofter refer to a specific allele, all equal hits SHOULD be annoted via this qualifier (which can be use multiple times). Also see the note on the limitations of the IMGT/GENE-DB reference database above.
- Ambiguous gene: Pick one, annotate using the qualifiers as noted for ambiguous allele.

JUNCTION annotation

INSDC does currently not define a key to annotate JUNCTION¹⁰. Therefore the following procedure MUST be used:

- REQUIRED: key CDS, indicating the positions of
 - 1. the first bp of the first AA of JUNCTION
 - 2. the last bp of the last AA of JUNCTION as determined by the utilized V(D)J inference tool.

Open coordinates MUST be used for both coordinates to allow for automated creation of the /translated qualifier providing the peptide sequence. Further note that a non-productive JUNCTION can have a length not divisible by three. This key contains the following qualifiers:

- REQUIRED: qualifier /codon_start with the assigned value "1".
- REQUIRED: qualifier / function with the assigned value "JUNCTION".
- REQUIRED: qualifier /product with an assigned value matching the regular expression

¹⁰ NCBI confirmed that once there would be enough datasets using the *JUNCTION* tag as specified here, a motion for an INSDC-sanctioned key could be initiated.

The variable region referred to in the string MUST be the same as the one given in the misc_feature key.

- RECOMMENDED: qualifier /inference, indicating the tool used for positional inference. The description string SHOULD use COORDINATES as category and protein motif as type [INSDC_FT].
- FORBIDDEN: qualifier /translated, which will be automatically added by Genbank.

Note that the complete CDS key will be removed by Genbank if the translation contains stop codons or to many "N" (exact number unknown). As such a record will lack a central piece of REQUIRED information it is RECOMMENDED that submitters either

- remove the complete record or
- replace the CDS with a misc_feature key while at the same time removing the /codon_start and /product qualifiers

upfront, as described in the submission manual. If the submitter chooses the replacement option, it has to be ensured that the annotated coordinates are actually valid and not affect by the frame- shift.

Record body

The record body starts after ORIGIN (ENA: SQ line) and MUST contain:

• the consensus sequence

References

Footnotes

Appendix

Example record (GenBank format)

LOCUS	AB123456 420 bp mRNA linear EST 01-JAN-2015
DEFINITION	TLS: Mus musculus immunoglobulin heavy chain variable region,
	sequence.
ACCESSION	AB123456
VERSION	AB123456.7
KEYWORDS	TLS; Targeted Locus Study; AIRR; MiAIRR:1.0.
SOURCE	Mus musculus
ORGANISM	Mus musculus
	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata;
	Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Glires;
	Rodentia; Sciurognathi; Muroidea; Muridae; Murinae; Mus.
REFERENCE	1 (bases 1 to 420)
AUTHORS	Stibbons,P.
TITLE	Section 5 information for experiment FOO1
JOURNAL	published (01-JAN-2000) on Zenodo
REMARK	DOI:10.1000/0000-12345678
REFERENCE	2 (bases 1 to 420)

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(continued from previous page)
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AUTHORS	Stibbons,P.					
TITLE	Direct Submission					
JOURNAL	Submitted (01-JAN-2000) Center for Transcendental Immunology,					
	Unseen University, Ankh-Morpork, 12345, DISCWORLD					
DBLINK	-	ct: PRJNA000001				
	BioSample	e: SAMN000001				
	Sequence	Read Archive: SRR0000001				
FEATURES		Location/Qualifiers				
source	!	1420				
		/organism="Mus musculus"				
		/mol_type="mRNA"				
		/strain="C57BL/6J"				
		/citation=[1]				
		/rearranged				
		/note="AIRR_READ_COUNT:123"				
V_regi		1324				
misc_f	eature	1324				
		<pre>/note="immunoglobulin heavy chain variable region"</pre>				
V_segm	lent	1257				
		/gene="IGHV1-34"				
		/allele="01"				
		/db_xref="IMGT/LIGM:AC073565"				
		<pre>/inference="COORDINATES:alignment:IgBLAST:1.6"</pre>				
D_segm	lent	266272				
		/gene="IGHD2-2"				
		/allele="01"				
		/db_xref="IMGT/LIGM:AJ851868"				
		<pre>/inference="COORDINATES:alignment:IgBLAST:1.6"</pre>				
J_segm	lent	291324				
		/gene="IGHJ4"				
		/allele="01"				
		/db_xref="IMGT/LIGM:V00770"				
~ ~ ~		<pre>/inference="COORDINATES:alignment:IgBLAST:1.6"</pre>				
CDS		<258>290				
		/codon_start=1				
		/function="JUNCTION"				
		<pre>/product="immunoglobulin heavy chain junction region" /i.c. "Boooppluating the state of a place of a state of a stat</pre>				
		/inference="COORDINATES:protein motif:IgBLAST:1.6"				
		/translated="CARAGVYDGYTMDYW"				
C_regi	on	325420				
ODICIN		/gene="Ighg2c"				
ORIGIN						
		ttcagtgaag atgtcctgca aggcttctgg ctacacattc actgactata				
		ggtgaagcag agccatggaa agagccttga gtggattgca tatattaatc				
		tggttatggc tataacgaca agttcaggga caaggccaca ttgactgtcg				
		caacacagec tacatgggge teegeageet gaeetetgag gaetetgeag				
		tgcaagagcg ggagtttacg acggatatac tatggactac tggggtcaag				
		cacegtetee teagecaaaa caacageeee ateggtetat ceaetggeee				
	lgtgtgtgg	aggtacaact ggctcctcgg tgactctagg atgcctggtc aagggcaact				
//						

Glossary

- MUST / REQUIRED: Indicates that an element or action is necessary to conform to the standard.
- SHOULD / RECOMMENDED: Indicates that an element or action is considered to be best practice by AIRR,

but not necessary to conform to the standard.

- CAN / OPTIONAL: Indicates that it is at the discretion of the user to use an element or perform an action.
- MUST NOT / FORBIDDEN: Indicates that an element or action will be in conflict with the standard.

Abbreviations

- AA: amino acid
- bp: base pair
- DOI: digital object identifier
- FT: INSDC Feature Table
- INSDC: International Nucleotide Sequence Database Collaboration
- SRA: sequence read archive

Introduction

The MiAIRR standard

The MiAIRR standard (minimal information about adaptive immune receptor repertoires) is a minimal reporting standard for experiments using sequencing-based technologies to study adaptive immune receptors (e.g. T cell receptors or immunoglobulins). It is developed and maintained by the Minimal Standards Working Group of the Adaptive Immune Receptors Repertoire (AIRR) Community [Breden_2017]. The current version (1.0) of the standard has been recently published [Rubelt_2017] and was passed by the general assembly at the annual AIRR Community meeting in December 2017. MiAIRR requires researchers to report six sets of information:

- 1. study, subject, diagnosis & intervention
- 2. sample collection
- 3. sample processing and sequencing
- 4. raw sequencing data
- 5. data processing
- 6. processed sequences with a basic analysis results

However, MiAIRR only describes the mandatory data items that have to be reported, but neither provides details how and where to deposit data nor specifies data types and formats. Therefore this document aims to provide both a submission manual for users as well as a detailed data specification for developers.

Requirement Levels of AIRR Schema Fields

Clarification of Terms

- The terms "MUST", "MUST NOT", "REQUIRED", "SHOULD", "SHOULD NOT", "RECOMMENDED", "MAY" and "OPTIONAL" are to be interpreted as described in [RFC2119].
- The terms "IF" and "ONLY IF" are are to be interpreted as sufficent and necessary requirement, respectively.

- The term "NULL-LIKE" is an extension of the NULL term in SQL and its equivalents in other programming languages, referring to the absence of data in a field (i.e., the field is empty). NULL-LIKE **additionally** includes the following terms, which also define the reason why the information is missing. As these terms are expected to be provided as text, the field would not be NULL but nevertheless NULL-LIKE (i.e., it lacks biologically interpretable information).
 - not_applicable: There is no meaningful value for this field due to study design (e.g., sex for a phage library).
 - not_collected: Data for this field was not collected during the study.
 - missing: Data for field was collected, but is not available now.

Categories of AIRR Schema Fields

- Fields MUST be indicated by the x-airr:miairr property IF and ONLY IF the field or its content is governed by the MiAIRR data standard [Rubelt_2017].
- The x-airr:miairr property MUST be assigned to one of the following three requirement levels:
 - essential: Information on this field MUST be provided and is considered critical for the meaningful interpretation of the data. Therefore the value of such a field MUST NOT be NULL-LIKE. Due to this strict requirement, this level is only assigned to a small set of fields. Importantly, fields are **not** elevated to this level based on the fact that the respective information should typically be available to the data generator. This was decided to simplify MiAIRR-compliant data annotation by third parties, who might perform this task based on publicly available information only.
 - important: Information for this field MUST be provided. However, the field MAY be assigned a NULL-LIKE value if the respective information is not available. The majority of fields governed by the MiAIRR data standard are assigned to this level.
 - defined: Information for this field MAY be provided. However, IF information matching the semantic definition of the field is provided, this field MUST be used for reporting.

Compliance with the MiAIRR Data Standard

- Compliance to the MiAIRR Data Standard is currently a binary state, i.e., a data either is or is not compliant, there are not "grades" of compliance. However, additional requirements for specific use cases might be defined in the future.
- Data sets are considered MiAIRR-compliant ONLY IF all essential and important fields are reported.
- Note that important fields with NULL-LIKE values MUST NOT be dropped from a data set.
- Implementors of data entry interfaces SHOULD NOT set the default value of important fields to NULL-LIKE values, i.e., users should be required to actively select the values.

Metadata Annotation Guidelines

Purpose of this Document

This document describes the RECOMMENDED ways to provide metadata annotation for various experimental setups.

Clarification of Terms

• The key words "MUST", "MUST NOT", "REQUIRED", "SHOULD", "SHOULD NOT", "RECOM-MENDED", "MAY", and "OPTIONAL" in this document are to be interpreted as described in [RFC2119].

Individual fields

library_generation_method

The library_generation_method describes how the nucleic acid annotated in template_class that encodes the V(D)J-rearrangement it reverse-transcribed, amplified and/or otherwise prepared for further processing. Typically this procedure will precede further NGS platform- specific steps, however these procedures MAY be combined. The field uses a controlled vocabulary, the individual values are described below:

template	e_dliabsmsary_genera	t Methodology
DNA	PCR	Conventional PCR on genomic DNA of a vertebrate host (requires:
		<pre>synthetic == false)</pre>
		Conventional PCR on DNA of a synthetic library (requires: synthetic ==
		true)
RNA	RT (RHP) +PCR	RT-PCR using random hexamer primers
	RT(oligo-dT)+P	CIRT-PCR using oligo-dT primers
	RT(oligo-dT)+T	S-5P-REACE PCR (i.e. RT is followed by a template switch (TS) step) using
		oligo-dT primers
	RT(oligo-dT)+T	S 5CHRAGE PER using oligo-dT primers and template switch primers contain-
		ing unique molecular identifiers (UMI), i.e., the 5' end is UMI-coded
	· · · · · · · · · · · · · · · · · · ·	CRT-PCR using transcript-specific primers
	· · · · · · · · · · · · · · · · · · ·	S-5P-REACE PCR using transcript- specific primers
	RT(specific)+T	S 5CHRAGE PER using transcript- specific primers and template switch primers
		containing UMIs
	RT(specific+UM	I REPORE using transcript-specific primers containing UMIs (i.e., the 3' end is
		UMI-coded)
	RT(specific+UM	1 54-RACE RCR using transcript- specific primers containing UMIs (i.e., the 3'
		end is UMI-coded)
	RT(specific)+T	S RT-based generation of dsDNA without subsequent PCR. This is used by
		RNA-seq kits.
any	other	Any methodology not covered above

Specific Use Cases and Experimental Setups

Synthetic libraries

In synthetic libraries (e.g. phage or yeast display), particles present genetically engineered constructs (e.g. scFv fusion receptors) on their surface. As this deviates substantially from other workflows, the following annotation SHOULD/MUST be used:

- In general, Subject should be interpreted as the initial library that undergoes a mutation/selection procedure.
- synthetic: MUST be set to true
- species: It is assumed that every synthetic library is derived from V and J genes that exist in some vertebrate species. This field SHOULD encode this species. Importantly, it MUST NOT encode the phage vector, the

bacterial host or the comparable biological component of the library system that constitutes the presenting particle.

- sample_type: SHOULD be NULL.
- single_cell: Only true if individual particles are isolated and sequenced. Note that colonies or plaques, even if containing genetically identical particles, *per se* do not match this definition and therefore MUST be annotated as false.
- cell_storage: SHOULD be used for non-cellular particles analogously.
- physical_linkage: For scFv constructs the hetero_prelinkeded term MUST be used. VHH (i.e. camelid) libraries SHOULD annotate none as there is only a single rearrangement envolved.

2.3.2 AIRR Data Representations

AIRR Data Representations are versioned specifications that consist of a file format and a well-defined schema. The schema is provided in a machine-readable YAML document that follows the OpenAPI v2.0 specification. The schema defines the data model, field names, data types, and encodings for AIRR standard objects. Strict typing enables interoperability and data sharing between different AIRR-seq analysis tools and repositories, and some fields use a controlled vocabulary or an ontology for value restriction. Specification extensions are utilized to define AIRR-specific attributes.

FAIR Principles

We desire AIRR standard objects to be FAIR (findable, accessible, interoperable and reusable) [Wilkinson_2016]:

- findable: by giving AIRR standard objects a globally unique identifier
- accessible: by providing an API where AIRR standard objects can be queried and downloaded
- interoperable: by defining a OpenAPI schema for the AIRR standard objects
- reusable: by linking the AIRR standard objects together into a standard formats

AIRR Data Model

The MiAIRR standard defines the minimal information for submission and publication of AIRR-seq datasets. The standard defines a set of data elements for this information and organizes them into six high-level sets.

- Study, Subject and Diagnosis
- Sample Collection
- Sample Processing and Sequencing
- Raw Sequences
- Data Processing
- Processed Sequences with Annotations

However beyond these sets, MiAIRR does not define any structure, data model or relationship between the data elements. This provides flexibility for the information to be stored in various database repositories but is problematic for interoperability and reusability of that information by computer programs. The AIRR Data Model overcomes these issues by defining a schema for the MiAIRR data elements, structuring them within schema objects, defining the relationship between those objects, and defining a file format.

Here are the primary schema objects of the AIRR Data Model:

Schema	Description
Object	
Study	Information about the experimental study design, including the title of the study, laboratory contact
	information, funding, and linked publications.
-	Information about the study cohorts and individual subjects, including species, sex, age, and ancestry.
Diagnos	allaformation about disease state(s), therapies, and study group membership (e.g., control versus disease).
Sample	Information about the origin and expected composition of the biological sample(s). This set aims to
	capture essential information about the collection of a sample, including its source (e.g., anatomical
	site), its provenance (provider), and the experimental condition (e.g., the time point during the course
	of a disease or treatment).
CellPro	deformation about the cell subset being profiled, as defined by the investigator, and the flow cytometry
	or other markers used to select the subset. Additional information includes the number of cells per
	sample and whether cells were prepared in bulk or captured as single cells.
Nucleio	An fortimation about queleic acid sample type (e.g., RNA versus DNA) and how immune-receptor gene
	rearrangements were amplified and sequenced (for example, RACE-PCR versus multiplex PCR, paired
~	PCR, and/or varying read length and sequencing chemistries).
Sequend	information about the sequencing run, such as the number of reads, read lengths, quality control pa-
	rameters, the sequencing kit and instrument(s) used, and run batch number. Also includes information
	about the raw data for the sequencing run (e.g., FASTQ files).
	deformation about the data processing to transform the raw sequencing data into Rearrangements.
Reperto	Geomposite object that combines the schema objects Study, Subject, Diagnosis, Sample,
	CellProcessing, NucleicAcidProcessing, SequencingRun, and DataProcessing.
	Each Repertoire has a unique identifier repertoire_id for linking with other data files, e.g.
	Rearrangements. Repertoires have their own schema and file format described <i>here</i> .
Rearrar	Anenotated sequences describing adaptive immune receptor chains. Rearrangements have their own
	schema and file format described <i>here</i> .

Relationship between Schema Objects

The MiAIRR categories are hierarchical, and includes information about the study, the subjects, the collected samples and how they are processed, details of the sequencing protocol, and information about the data analysis. The top-down relationships are either 1-to-n indicating the top level object can be related to any number of sub-level objects, or n-to-n indicating any number of top level object can be related to any number of sub-level objects. Lastly, 1-to-1 indicates the top level object is related to a single sub-level object.

- Study 1-to-n with Subject. A study may contain any number of subjects.
- Subject 1-to-n with Diagnosis. Each subject may contain any number of diagnoses.
- Subject 1-to-n with Sample. Each subject may contain any number of samples.
- Sample 1-to-n with CellProcessing. A sample may have any number of cell processing records.
- CellProcessing 1-to-n with NucleicAcidProcessing. A cell processing record may have any number of nucleic acid processing records.
- NucleicAcidProcessing 1-to-n with SequencingRun. A nucleic acid processing records may have any number of sequencing runs.
- SequencingRun n-to-n with DataProcessing. Multiple sequencing runs can be combined in a data processing, and multiple data processing can be done on a sequencing run.

However, this hierarchy is deep and complicated. Therefore to simplify the processing of this information, we denormalized the hierarchy around the conceptual Repertoire object. This denormalization represents many relationships as 1-to-1 which simplifies the structure. A single Repertoire has these relationships with the primary schema objects.

- Repertoire 1-to-1 with Study. A repertoire is for a single study, though a study may have multiple repertoires.
- Repertoire 1-to-1 with Subject. A repertoire is for a single subject, though a subject may have other repertoires defined.
- Sample 1-to-1 with CellProcessing, NucleicAcidProcessing, and SequencingRun. A sample is associated with a single chain of sample processing from initial collection, through cell and nucleic acid processing, to sequencing.
- Repertoire 1-to-n with Sample. Generally a repertoire has a single sample, but sometimes studies perform technical replicates or re-sequencing to generate additional data, and these studies will have multiple samples, which are to be combined and analyzed together as part of the same repertoire.
- Repertoire 1-to-n with DataProcessing. A repertoire can be analyzed multiple times. More details about multiple data processing is provided below.

The trade-off with denormalization of the hierarchy is that it causes duplication of data. For example, two repertoires for the same study will have the Study information duplicated within each of the two repertoire records; likewise multiple repertoires for the same subject will have the Subject information duplicated.

While the denormalized Repertoire simplifies read-only access to the MiAIRR information, it complicates data entry and write access to the information because updates need to be propagated to all of the duplicate records. Therefore, Repertoire was designed to be easily transformed into a normalized form, representing the full hierarchy of the objects, by utilizing the *study_id*, *subject_id*, and *sample_id* fields to uniquely identify the Study, Subject and Sample objects across multiple repertoires. The exception is that CellProcessing and NucleicAcidProcessing do not have their own unique identifiers, so they are included within Sample.

AIRR extension properties

The OpenAPI V2 specification provides the ability to define extension properties on schema objects. These are additional properties on the schema definition directly, not to be confused with additional properties on the data. These extension properties allow those schema definitions to be annotated with MiAIRR and AIRR specific information. Instead of creating separate extensions for each property, a single extension x-airr property is defined, which is an object that contains any number of properties. Within the AIRR schema, AIRR_Extension defines the schema for the x-airr object and the properties within it. Here is a list of the currently supported AIRR extension properties:

Exten-	Description
sion	
miairr	Present if the annotated property is a MiAIRR data standard element. Always has a requirement level
	assigned to it.
nullabl	eAssumes miairr. False if the annotated property must not be NULL by the MiAIRR standard, other-
	wise True or null.
set	Assumes miairr. The MiAIRR set for the annotated property.
subset	Assumes miairr. The MiAIRR subset for the annotated property.
name	Assumes miairr. The MiAIRR field name.
format	Describes the format for the annotated property. Value is either free text, controlled
	vocabulary or ontology.
ontolog	yIf format=ontology then this provides additional information about the ontology including draft
	status, name, URL and top node term.

Schema Definitions

Repertoire Schema

A Repertoire is an abstract organizational unit of analysis that is defined by the researcher and consists of study metadata, subject metadata, sample metadata, cell processing metadata, nucleic acid processing metadata, sequencing run metadata, a set of raw sequence files, data processing metadata, and a set of Rearrangements. A Repertoire gathers all of this information together into a composite object, which can be easily accessed by computer programs for data entry, analysis and visualization.

A Repertoire is specific to a single subject otherwise it can consist of any number of samples (which can be processed in different ways), any number of raw sequence files, and any number of rearrangements. It can also consist of any number of data processing metadata objects that describe the processing of raw sequence files into Rearrangements.

Typically, a Repertoire corresponds to the biological concept of the immune repertoire for that single subject which the researcher experimentally measures and computationally analyzes. However, researchers can have different interpretations about what constitutes the biological immune repertoire; therefore, the Repertoire schema attempts to be flexible and broadly useful for all AIRR-seq studies.

Another researcher can take the same raw sequencing data and associated metadata and create their own Repertoire that is different from the original researcher's. A common example is to define a repertoire that is a subset such as "productive rearrangements for IGHV4" whereas the original researcher defined a more generic "B cell repertoire". This new Repertoire would have much of the same metadata as the original Repertoire, except associated with a different study, and with additional information in the data processing metadata that describes how the rearrangements were filtered down to just the "productive rearrangements for IGHV4". Likewise, another researcher may get access to the original biosample material and perform their own sample processing and sequencing, which also would be a new Repertoire. That new Repertoire could combine samples from the original researcher's Repertoire with the new sample data as a large dataset for the subject.

Multiple Data Processing on a Repertoire

Data processing can be a complicated multi-stage process. Documenting the process in a formal way is challenging because of the diversity of actions that may be performed. The MiAIRR standard requires documentation of the process but in an informal way with free text descriptions. A Repertoire might undergo multiple different data processing for any number of reasons, e.g. to compare the results from different toolchains, or to compare different settings for the same toolchain.

It is expected that all of the Samples of a Repertoire will be processed together within a DataProcessing. That is, a DataProcessing that only uses some but not all samples in a Repertoire could be confusing to users and appear as though data is missing. Likewise, processing some samples within a Repertoire with one DataProcessing and the remaining samples with a different DataProcessing could also confuse users. Because DataProcessing is unstructured information, it is not possible to validate that all Samples in a Repertoire are being processed together, so this expectation cannot be strictly enforced.

Having multiple DataProcessing for a Repertoire will create multiple sets of Rearrangements that are distinct and separate from each other. Analysis tools need to be careful not to mix these sets of Rearrangements from different DataProcessing because it can generate incorrect results. The identifier data_processing_id was added so Rearrangements can identify their specific DataProcessing.

Linking Data

Each Repertoire has a unique repertoire_id identifier. This identifier should be globally unique so that repertoires from multiple studies can be combined together without conflict. The repertoire_id is used to link other AIRR data to a Repertoire. Specifically, the *Rearrangements Schema* includes repertoire_id for referencing the specific Repertoire for that Rearrangement. If a Repertoire has multiple DataProcessing then data_processing_id should be used to distinguish the appropriate DataProcessing within the Repertoire. The Rearrangements contains data_processing_id for this purpose. The data_processing_id is only unique within a Repertoire so repertoire_id should first be used to get the appropriate Repertoire object and then data_processing_id used to acquire the appropriate DataProcessing.

It is expected that typical Repertoires might only have a single DataProcessing, in which case repertoire_id and data_processing_id will be semantically equivalent and only the former should be used.

If a Repertoire has multiple sample processing objects in the sample array then sample_processing_id should be used to distinguish the the appropriate sample processing object within the Repertoire. The Rearrangement object can contain a sample_processing_id to uniquely identify a sample processing object within a Repertoire. Like data_processing_id, the sample_processing_id is only unique within the Repertoire so repertoire_id should first be used to get the appropriate Repertoire object and then sample_processing_id should be used to determine the appropriate sample processing object that is associated with the Rearrangement. If the Rearrangement object does not have a sample_processing_id then it can be assumed that the rearrangement is associated with all of the samples in the Repertoire (e.g. the rearrangement is a collapsed rearrangement across multiple samples).

It is expected that Repertoires might often have a single sample processing object, in which case repertoire_id and sample_processing_id will be semantically equivalent and only the former should be used.

Finally, if it is necessary to link a Rearrangement object with a unique pairing of sample processing and DataProcessing, the repertoire_id of the Rearrangement object should be used to identify the correct Repertoire object and then the data_processing_id should be used to identify the correct DataProcessing metadata and the sample_processing_id should be used to identify the correct sample processing metadata within that Repertoire.

Duality between Repertoires and Rearrangements

There is an important duality relationship between Repertoires and Rearrangements, specifically with the experimental protocols described in the Repertoire versus the annotations on Rearrangements. A Repertoire defines an experimental design for what a researcher intends to measure or observe, while the Rearrangements are what was actually measured and observed. Technically, the border between the two occurs at sequencing, that is when the biological physical entity (prepared DNA) is measured and recorded as information (nucleotide sequence).

This duality is important when considering how to answer certain questions. For example, locus for Rearrangements may have the value "IGH" which indicates that B cell heavy chain receptors were measured, yet the Repertoire might have "T cell" in cell_subset which indicates the researcher intended to measure T cells. This conflict between the two indicates something is wrong. Differences can occur in many ways, as with errors in the experimental protocol, or data processing might have incorrectly processed the raw sequencing data leading to invalid annotations.

File Format Specification

Files are YAML/JSON with a structure defined below. Files should be encoded as UTF-8. Identifiers are case-sensitive. Files should have the extension .yaml, .yml, or .json.

File Structure

• The file as a whole is considered a dictionary (key/value pair) structure with the keys Info and Repertoire.

- The file can (optionally) contain an Info object, at the beginning of the file, based upon the Info schema in the OpenAPI V2 specification. If provided, version in Info should reference the version of the AIRR schema for the file.
- The file should correspond to a list of Repertoire objects, using Repertoire as the key to the list.
- Each Repertoire object should contain a top-level key/value pair for repertoire_id that uniquely identifies the repertoire.
- Some fields require the use of a particular ontology or controlled vocabulary.
- The structure is the same regardless of whether the data is stored in a file or a data repository. For example, The *ADC API* will return a properly structured JSON object that can be saved to a file and used directly without modification.

Repertoire Fields

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Name	Туре	Attributes	Definition
repertoire_id	string	optional, iden-	Identifier for the repertoire object. This identifier should
		tifier, nullable	be globally unique so that repertoires from multiple
			studies can be combined together without conflict. The
			repertoire_id is used to link other AIRR data to a Reper-
			toire. Specifically, the Rearrangements Schema in-
			cludes repertoire_id for referencing the specific Reper-
			toire for that Rearrangement.
repertoire_name	e string	optional, nul-	Short generic display name for the repertoire
		lable	
repertoire_desc	r strpntg ion	optional, nul-	Generic repertoire description
		lable	
study	Study	required	Study object
subject	Subject	required	Subject object
sample	array	required	List of Sample objects
data_processing	f array of <i>Dat</i> -	required	List of Data Processing objects
	aProcessing		

Study Fields

Name	Туре	Attributes	Definition
study_id	string	required, nul- lable	Unique ID assigned by study registry
study_title	string	required, nul- lable	Descriptive study title
study_type	Ontology	required, nul- lable	Type of study design
study_descripti	atring	optional, nul- lable	Generic study description
inclusion_exclu	ıs ströng_ criter:	arequired, nul- lable	List of criteria for inclusion/exclusion for the study
grants	string	required, nul- lable	Funding agencies and grant numbers
collected_by	string	required, nul- lable	Full contact information of the data collector, i.e. the person who is legally responsible for data collection and release. This should include an e-mail address.
lab_name	string	required, nul- lable	Department of data collector
lab_address	string	required, nul- lable	Institution and institutional address of data collector
submitted_by	string	required, nul- lable	Full contact information of the data depositor, i.e. the person submitting the data to a repository. This is sup- posed to be a short-lived and technical role until the sub- mission is relased.
pub_ids	string	required, nul- lable	Publications describing the rationale and/or outcome of the study
keywords_study	array of string	required, nul- lable	Keywords describing properties of one or more data sets in a study

Subject Fields

Name	Туре	Attributes	Definition
subject_id	string	required, nul- lable	Subject ID assigned by submitter, unique within study
synthetic	boolean	required	TRUE for libraries in which the diversity has been syn- thetically generated (e.g. phage display)
species	Ontology	required	Binomial designation of subject's species
organism	Ontology	DEPRECATED	Binomial designation of subject's species
sex	string	required, nul- lable	Biological sex of subject
age_min	number	required, nul- lable	Specific age or lower boundary of age range.
age_max	number	required, nul- lable	Upper boundary of age range or equal to age_min for specific age. This field should only be null if age_min is null.
age_unit	Ontology	required, nul- lable	Unit of age range
age_event	string	required, nul- lable	Event in the study schedule to which <i>Age</i> refers. For NCBI BioSample this MUST be <i>sampling</i> . For other implementations submitters need to be aware that there is currently no mechanism to encode to potential delta between <i>Age event</i> and <i>Sample collection time</i> , hence the chosen events should be in temporal proximity.
age	string	DEPRECATED	
ancestry_popula	tströng	required, nul- lable	Broad geographic origin of ancestry (continent)
ethnicity	string	required, nul- lable	Ethnic group of subject (defined as cultural/language- based membership)
race	string	required, nul- lable	Racial group of subject (as defined by NIH)
strain_name	string	required, nul- lable	Non-human designation of the strain or breed of animal used
linked_subjects	s string	required, nul- lable	Subject ID to which Relation type refers
link_type	string	required, nul- lable	Relation between subject and <i>linked_subjects</i> , can be genetic or environmental (e.g.exposure)
diagnosis	array of <i>Diag-</i> nosis	optional	Diagnosis information for subject

Diagnosis Fields

Name	Туре	Attributes	Definition
study_group_des	scstringtion	required, nul-	Designation of study arm to which the subject is as-
		lable	signed to
disease_diagnos	i Q ntology	required, nul-	Diagnosis of subject
		lable	
disease_length	string	required, nul-	Time duration between initial diagnosis and current in-
		lable	tervention
disease_stage	string	required, nul-	Stage of disease at current intervention
		lable	
prior_therapies	string	required, nul-	List of all relevant previous therapies applied to subject
		lable	for treatment of Diagnosis
immunogen	string	required, nul-	Antigen, vaccine or drug applied to subject at this inter-
		lable	vention
intervention	string	required, nul-	Description of intervention
		lable	
medical_history	⁷ string	required, nul-	Medical history of subject that is relevant to assess the
		lable	course of disease and/or treatment

Sample Fields

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Name	Туре	Attributes	Definition
sample_id	string	required, nul-	Sample ID assigned by submitter, unique within study
		lable	
sample_type	string	required, nul-	The way the sample was obtained, e.g. fine-needle aspi-
		lable	rate, organ harvest, peripheral venous puncture
tissue	Ontology	required, nul-	The actual tissue sampled, e.g. lymph node, liver, pe-
		lable	ripheral blood
anatomic_site	string	required, nul-	The anatomic location of the tissue, e.g. Inguinal, femur
		lable	
disease_state_s	attionle	required, nul-	Histopathologic evaluation of the sample
		lable	
collection_time	_ stroinig nt_relat	required, nul-	Time point at which sample was taken, relative to Col-
		lable	lection time event
collection_time	_ stroinig nt_refe	erequired, nul-	Event in the study schedule to which Sample collection
		lable	time relates to
biomaterial_pro	vströhegr	required, nul-	Name and address of the entity providing the sample
		lable	

Tissue and Cell Processing Fields

Name	Туре	Attributes	Definition
tissue_processi	rstgring	required, nul-	Enzymatic digestion and/or physical methods used to
		lable	isolate cells from sample
cell_subset	Ontology	required, nul- lable	Commonly-used designation of isolated cell population
cell_phenotype	string	required, nul- lable	List of cellular markers and their expression levels used to isolate the cell population
cell_species	Ontology	optional, nul- lable	Binomial designation of the species from which the an- alyzed cells originate. Typically, this value should be identical to <i>species</i> , if which case it SHOULD NOT be set explicitly. Howver, there are valid experimental se- tups in which the two might differ, e.g. chimeric animal models. If set, this key will overwrite the <i>species</i> infor- mation for all lower layers of the schema.
single_cell	boolean	required, nul- lable	TRUE if single cells were isolated into separate com- partments
cell_number	integer	required, nul- lable	Total number of cells that went into the experiment
cells_per_react	iioteger	required, nul- lable	Number of cells for each biological replicate
cell_storage	boolean	required, nul- lable	TRUE if cells were cryo-preserved between isolation and further processing
cell_quality	string	required, nul- lable	Relative amount of viable cells after preparation and (if applicable) thawing
cell_isolation	string	required, nul- lable	Description of the procedure used for marker-based iso- lation or enrich cells
cell_processing	<u>s</u> tring ocol	required, nul- lable	Description of the methods applied to the sample in- cluding cell preparation/ isolation/enrichment and nu- cleic acid extraction. This should closely mirror the Ma- terials and methods section in the manuscript.

Nucleic Acid Processing Fields

Name	Туре	Attributes	Definition
template_class	string	required	The class of nucleic acid that was used as primary start-
			ing material for the following procedures
template_qualit	ystring	required, nul-	Description and results of the quality control performed
		lable	on the template material
template_amount	string	required, nul-	Amount of template that went into the process
		lable	
library_generat	-	required	Generic type of library generation
library_generat	i stmi<u>ng</u>protoco l		Description of processes applied to substrate to obtain a
		lable	library that is ready for sequencing
library_generat	i stmi<u>n</u>g kit_ver:		When using a library generation protocol from a com-
		lable	mercial provider, provide the protocol version number
pcr_target	array of <i>PCR</i> -	optional	If a PCR step was performed that specifically targets the
	Target		IG/TR loci, the target and primer locations need to be
			provided here. This field holds an array of PCRTarget
			objects, so that multiplex PCR setups amplifying mul-
			tiple loci at the same time can be annotated using one
			record per locus. PCR setups not targeting any specific
			locus must not annotate this field but select the appro- priate library_generation_method instead.
	- cotmin a	raquirad	
complete_sequer	i Cseising	required	To be considered <i>complete</i> , the procedure used for library construction MUST generate sequences that 1) in-
			clude the first V gene codon that encodes the mature
			polypeptide chain (i.e. after the leader sequence) and
			2) include the last complete codon of the J gene (i.e. 1
			bp 5' of the J->C splice site) and 3) provide sequence
			information for all positions between 1) and 2). To be
			considered <i>complete & untemplated</i> , the sections of the
			sequences defined in points 1) to 3) of the previous sen-
			tence MUST be untemplated, i.e. MUST NOT over-
			lap with the primers used in library preparation. <i>mixed</i>
			should only be used if the procedure used for library
			construction will likely produce multiple categories of
			sequences in the given experiment. It SHOULD NOT
			be used as a replacement of a NULL value.
physical_linkag	estring	required	In case an experimental setup is used that physically
			links nucleic acids derived from distinct Rearrange-
			ments before library preparation, this field describes the
			mode of that linkage. All <i>hetero_*</i> terms indicate that
			in case of paired-read sequencing, the two reads should
			be expected to map to distinct IG/TR loci. *_head-head
			refers to techniques that link the 5' ends of transcripts
			in a single-cell context. *_ <i>tail-head</i> refers to techniques
			that link the 3' end of one transcript to the 5' end of an-
			other one in a single-cell context. This term does not
			provide any information whether a continuous reading-
			frame between the two is generated. *_ <i>prelinked</i> refers
			to constructs in which the linkage was already present
			on the DNA level (e.g. scFv).

PCR Target Locus Fields

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Name	Туре	Attributes	Definition
pcr_target_locu	sstring	required, nul-	Designation of the target locus. Note that this field
		lable	uses a controlled vocubulary that is meant to provide
			a generic classification of the locus, not necessarily the
			correct designation according to a specific nomencla-
			ture.
forward_pcr_pri	m stning target_i	l vequired , nul-	Position of the most distal nucleotide templated by the
		lable	forward primer or primer mix
reverse_pcr_pri	m stning target_i	l ocquired n nul-	Position of the most proximal nucleotide templated by
		lable	the reverse primer or primer mix

Raw Sequence Data Fields

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Name	Туре	Attributes	Definition
file_type	string	required, nul-	File format for the raw reads or sequences
		lable	
filename	string	required, nul-	File name for the raw reads or sequences. The first file
		lable	in paired-read sequencing.
read_direction	string	required, nul-	Read direction for the raw reads or sequences. The first
		lable	file in paired-read sequencing.
read_length	integer	required, nul-	Read length in bases for the first file in paired-read se-
		lable	quencing
paired_filename	e string	required, nul-	File name for the second file in paired-read sequencing
		lable	
paired_read_dir	estatinigon	required, nul-	Read direction for the second file in paired-read se-
		lable	quencing
paired_read_ler	ginteger	required, nul-	Read length in bases for the second file in paired-read
		lable	sequencing

Sequencing Run Fields

Name	Туре	Attributes	Definition
sequencing_run_	istring	required, nul-	ID of sequencing run assigned by the sequencing facility
		lable	
total_reads_pas	sintegerqc_filt	æquired, nul-	Number of usable reads for analysis
		lable	
sequencing_plat	fstning	required, nul-	Designation of sequencing instrument used
		lable	
sequencing_faci	lstring	required, nul-	Name and address of sequencing facility
		lable	
sequencing_run_	chartineg	required, nul-	Date of sequencing run
		lable	
sequencing_kit	string	required, nul-	Name, manufacturer, order and lot numbers of sequenc-
		lable	ing kit
sequencing_file	sRawSequenceD	atoptional	Set of sequencing files produced by the sequencing run

Data Processing Fields

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Name	Туре	Attributes	Definition
data_processing	<u>ströng</u>	optional, iden-	Identifier for the data processing object.
		tifier, nullable	
primary_annotat	iboolean	optional, iden-	If true, indicates this is the primary or default data pro-
		tifier	cessing for the repertoire and its rearrangements. If
			false, indicates this is a secondary or additional data pro-
			cessing.
software_version	rstring	required, nul-	Version number and / or date, include company
		lable	pipelines
paired_reads_as	sseminingly	required, nul-	How paired end reads were assembled into a single re-
		lable	ceptor sequence
quality_thresho	lstrising	required, nul-	How sequences were removed from (4) based on base
		lable	quality scores
primer_match_cu	tstnings	required, nul-	How primers were identified in the sequences, were they
		lable	removed/masked/etc?
collapsing_meth	ostring	required, nul-	The method used for combining multiple sequences
		lable	from (4) into a single sequence in (5)
data_processing	<u>s</u> tningtocols	required, nul-	General description of how QC is performed
		lable	
data_processing	_afrialyeosf string	optional, nul-	Array of file names for data produced by this data pro-
		lable	cessing.
germline_databa	ssering	required, nul-	Source of germline V(D)J genes with version number or
		lable	date accessed.
analysis_prover	astring_id	optional, nul-	Identifier for machine-readable PROV model of analysis
		lable	provenance

Rearrangement Schema

A Rearrangement is a sequence which describes a rearranged adaptive immune receptor chain (e.g., antibody heavy chain or TCR beta chain) along with a host of annotations. These annotations are defined by the AIRR Rearrangement schema and comprises eight categories.

Cate-	Description
gory	
Input	The input sequence to the V(D)J assignment process.
Identi-	Primary and foreign key identifiers for linking AIRR data across files and databases.
fiers	
Primary	The primary outputs of the V(D)J assignment process, which includes the gene locus, V, D, J, and
Annota-	C gene calls, various flags, V(D)J junction sequence, copy number (duplicate_count), and the
tions	number of reads contributing to a consensus input sequence (consensus_count).
Align-	Detailed alignment annotations including the input and germline sequences used in the alignment;
ment	score, identity, statistical support (E-value, likelihood, etc); and the alignment itself through CIGAR
Annota-	strings for each aligned gene.
tions	
Align-	The start/end positions for genes in both the input and germline sequences.
ment	
Posi-	
tions	
Region	Sequence annotations for the framework regions (FWRs) and complementarity-determining regions
Se-	(CDRs).
quence	
Region	Positional annotations for the framework regions (FWRs) and complementarity-determining regions
Posi-	(CDRs).
tions	
Junction	Lengths for junction sub-regions associated with aspects of the V(D)J recombination process.
Lengths	

File Format Specification

Data for Rearrangement or Alignment objects are stored as rows in a *tab-delimited* file and should be compatible with any TSV reader. A dataset is defined in this context as: a TSV file, a TSV with a companion YAML file containing metadata, or a directory containing multiple TSV files and YAML files.

Encoding

- The file should be encoded as ASCII or UTF-8.
- Everything is case-sensitive.

Dialect

- The record separator is a newline n and the field separator is a tab t.
- Fields or data should not be quoted.
- A header line with the AIRR-specified column names is always required.
- Values must not contain tab or newline characters.
- Values should avoid @, #, and quote (" or ') characters, as the result may be implementation dependent.
- Nested delimiters are not supported by the schema explicitly and should be avoided. However, if multiple values must be reported in a single column for an application specific reason, then the use of a comma as the delimiter is recommended.

File names

AIRR formatted TSV files should end with .tsv.

File Structure

The data file has two sections in this order:

- 1. Header. A single line with column names.
- 2. Data values. One record per line.

A comment section preceding the header (e.g., # or @ blocks) is not part of the specification, but such a section is reserved for potential inclusion in a future release. As such, a comment section should not be included in the file as it *may* be incompatible with a future specification.

Header

A single line containing the column names and specifying the field order. Any field that corresponds to one of the defined fields should use the specified field name.

Required columns

Some of the fields are defined as required and therefore must always be present in the header. Note, however, that all columns allow for null values. Therefore, required columns exist to define a core set of fields that are always present in the table structure, but do not mandate that a value be reported.

Custom columns

There are no restrictions on inclusion of additional custom columns in the Rearrangements file, provided such columns do not use the same name as an existing required or optional field. It is recommended that custom fields follow the same naming scheme as existing fields. Meaning, snake_case with narrowing scope when read from left to right. For example, sequence_id is the *"identifier* of the *query sequence"*.

Consider submitting a pull request for a field name reservation to the airr-standards repository if the field may be broadly useful.

Ordering

There are no requirements that fields or records be sorted or ordered in any specific way. However, the field ordering provided by the schema is a recommended default, with top-to-bottom equating to left-to-right.

Data Values

The possible data types are string, boolean, number (floating point), integer, and null (empty string).

Boolean values

Boolean values must be encoded as ${\mathbb T}$ for true and ${\mathbb F}$ for false.

Null values

All fields may contain null values. This includes columns that are described as required. A null value should be encoded as an empty string.

Coordinate numbering

All alignment sequence coordinates use the same scheme as IMGT and INSDC (DDBJ, ENA, GenBank), with the exception that partial coordinate information should not be used in favor of simply assigning the start/end of the alignment. Meaning, coordinates should be provided as 1-based values with closed intervals, without the use of > or < annotations that denoted a partial region.

CIGAR specification

Alignments details are specified using the CIGAR format as defined in the SAM specifications, with some vocabulary restrictions on the use of clipping, skipping, and padding operators.

The CIGAR string defines the reference sequence as the germline sequence of the given gene or region; e.g., for v_cigar the reference is the V gene germline sequence. The query sequence is what was input into the alignment tool, which must correspond to what is contained in the sequence field of the Rearrangement data. For the majority of use cases, this will necessarily exclude alignment spacers from the CIGAR string, such as IMGT numbering gaps. However, any gaps appearing in the query sequence should be accounted for in the CIGAR string so that the alignment between the query and reference is correctly represented.

The valid operator sets and definitions are as follows:

Operator	Description
=	An identical non-gap character.
Х	A differing non-gap character.
М	A positional match in the alignment. This can be either an identical (=) or differing (x)
	non-gap character.
D	Deletion in the query (gap in the query).
Ι	Insertion in the query (gap in the reference).
S	Positions that appear in the query, but not the reference. Used exclusively to denote the start
	position of the alignment in the query. Should precede any N operators.
N	A space in the alignment. Used exclusively to denote the start position of the alignment in
	the reference. Should follow any S operators.

Note, the use of either the =/X or M syntax is valid, but should be used consistently. While leading S and N operators are required, tailing S and N operators are optional.

For example, an D gene alignment that starts at position 419 in the query sequence (leading 418S), that is 16 nucleotides long with no indels (middle 16M), has an 10 nucleotide 5' deletion (leading 10N), a 5 nucleotide 3' deletion (trailing 5N), and ends 72 nucleotides from the end of the query sequence (trailing 71S) would have the following D gene CIGAR string (d_cigar) and positional information:

Field	Value
d_cigar	418S10N16M71S5N
d_sequence_start	419
d_sequence_end	434
d_germline_start	11
d_germline_end	26

Definition Clarifications

Junction versus CDR3

We work with the IMGT definitions of the junction and CDR3 regions. Specifically, the IMGT JUNCTION includes the conserved cysteine and tryptophan/phenylalanine residues, while CDR3 excludes those two residues. Therefore, our junction_aa fields which represent the extracted sequence include the two conserved residues, while the coordinate fields (cdr3_start and cdr3_end) exclude them.

Productive

The schema does not define a strict definition of a productive rearrangement. However, the IMGT definition is recommended:

- 1. Coding region has an open reading frame
- 2. No defect in the start codon, splicing sites or regulatory elements.
- 3. No internal stop codons.
- 4. An in-frame junction region.

Locus names

A naming convention for locus names is not strictly enforced, but the IMGT locus names are recommended. For example, in the case of human data, this would be the set: IGH, IGK, IGL, TRA, TRB, TRD, or TRG.

Gene and allele names

Gene call examples use the IMGT nomenclature, but no specific gene or allele nomenclature is strictly mandated. Species denotations may or may not be included in the gene name, as appropriate. For example, "Homo sapiens IGHV4-59*01", "IGHV4-59*01" and "AB019438" are all valid entries for the same allele.

However, when using an established reference database to assign gene calls adherence to the exact nomenclature used by the reference database is strongly recommended, as this will facilitate mapping to the database entries, cross-study comparison, and upload to public repositories.

Alignments

There is no required alignment scheme for the nucleotide and amino acid alignment fields. These fields may, or may not, include numbering spacers (e.g., IMGT-numbering gaps), variations in case to denote mismatches, deletions, or other features appropriate to the tool that performed the alignment. The only strict requirement is that the query ("sequence") and reference ("germline") **must** be properly aligned.

Fields

The specification includes two classes of fields. Those that are required and those that are optional. Required is defined as a column that must be present in the header of the TSV. Optional is defined as column that may, or may not, appear in the TSV. All fields, including required fields, are nullable by assigning an empty string as the value. There are no requirements for column ordering in the schema, although the Python and R reference APIs enforce ordering for the sake of generating predictable output. The set of optional fields that provide alignment and region coordinates ("_start"

and "_end" fields) are defined as 1- based closed intervals, similar to the SAM, VCF, GFF, IMGT, and INDSC formats (GenBank, ENA, and DDJB; http://www.insdc.org).

Most fields have strict definitions for the values that they contain. However, some commonly provided information cannot be standardized across diverse toolchains, so a small selection of fields have context-dependent definitions. In particular, these context-dependent fields include the optional "_score," "_identity," and "_support" fields used for assessing the quality of alignments which vary considerably in definition based on the methodology used. Similarly, the "_alignment" fields require strict alignment between the corresponding observed and germline sequences, but the manner in which that alignment is conveyed is somewhat flexible in that it allows for any numbering scheme (e.g., IMGT or KABAT) or lack thereof.

By default, data elements representing sequences in the schema contain nucleotide sequences except for data elements ending in "_aa," which are amino acid translations of the associated nucleotide sequence.

While the format contains an extensive list of reserved field names, there are no restrictions on inclusion of custom fields in the TSV file, provided such custom fields have a unique name. Furthermore, suggestions for extending the format with additional reserved names are welcomed through the issue tracker on the GitHub repository (https: //github.com/airr-community/airr-standards).

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Name	Туре	Attributes	Definition
sequence_id	string	required, identifier, nullable	Unique query sequence identifier for the Rearrangment. Most often this will be the input sequence header or a substring thereof, but may also be a custom identi- fier defined by the tool in cases where query sequences have been combined in some fashion prior to align- ment. When downloaded from an AIRR Data Commons repository, this will usually be a universally unique record locator for linking with other objects in the AIRR Data Model.
sequence	string	required, nul- lable	The query nucleotide sequence. Usually, this is the un- modified input sequence, which may be reverse comple- mented if necessary. In some cases, this field may con- tain consensus sequences or other types of collapsed in- put sequences if these steps are performed prior to align- ment.
sequence_aa	string	optional, nul- lable	Amino acid translation of the query nucleotide se- quence.
rev_comp	boolean	required, nul- lable	True if the alignment is on the opposite strand (reverse complemented) with respect to the query sequence. If True then all output data, such as alignment coordinates and sequences, are based on the reverse complement of 'sequence'.
productive	boolean	required, nul- lable	True if the V(D)J sequence is predicted to be productive.
vj_in_frame	boolean	optional, nul- lable	True if the V and J gene alignments are in-frame.
stop_codon	boolean	optional, nul- lable	True if the aligned sequence contains a stop codon.

Name	Туре	Attributes	Definition
complete_vdj	boolean	optional, nul-	True if the sequence alignment spans the entire V(D)J
comprete_vaj	boolean	lable	region. Meaning, sequence_alignment includes both the
			first V gene codon that encodes the mature polypeptide
			chain (i.e., after the leader sequence) and the last com-
			plete codon of the J gene (i.e., before the J-C splice site).
			This does not require an absence of deletions within the
			internal FWR and CDR regions of the alignment.
1	atrina	antional nul	
locus	string	optional, nul- lable	Gene locus (chain type). Note that this field uses a con-
		lable	trolled vocabulary that is meant to provide a generic
			classification of the locus, not necessarily the correct
			designation according to a specific nomenclature.
v_call	string	required, nul-	V gene with allele. If referring to a known reference
		lable	sequence in a database the relevant gene/allele nomen-
			clature should be followed (e.g., IGHV4-59*01 if using
			IMGT/GENE-DB).
d_call	string	required, nul-	First or only D gene with allele. If referring to a known
		lable	reference sequence in a database the relevant gene/allele
			nomenclature should be followed (e.g., IGHD3-10*01 if
			using IMGT/GENE-DB).
d2_call	string	optional, nul-	Second D gene with allele. If referring to a known ref-
		lable	erence sequence in a database the relevant gene/allele
			nomenclature should be followed (e.g., IGHD3-10*01
			if using IMGT/GENE-DB).
j_call	string	required, nul-	J gene with allele. If referring to a known reference
		lable	sequence in a database the relevant gene/allele nomen-
			clature should be followed (e.g., IGHJ4*02 if using
			IMGT/GENE-DB).
c_call	string	optional, nul-	Constant region gene with allele. If referring to a known
		lable	reference sequence in a database the relevant gene/allele
			nomenclature should be followed (e.g., IGHG1*01 if
			using IMGT/GENE-DB).
sequence_alignr	nestating	required, nul-	Aligned portion of query sequence, including any indel
		lable	corrections or numbering spacers, such as IMGT-gaps.
			Typically, this will include only the V(D)J region, but
			that is not a requirement.
sequence_alignr	neatutinga a	optional, nul-	Amino acid translation of the aligned query sequence.
		lable	
germline_alignr	neatuting	required, nul-	Assembled, aligned, full-length inferred germline
		lable	sequence spanning the same region as the se-
			quence_alignment field (typically the V(D)J region) and
			including the same set of corrections and spacers (if
			any).
germline_alignr	neatutinga a	optional, nul-	Amino acid translation of the assembled germline se-
		lable	quence.
junction	string	required, nul-	Junction region nucleotide sequence, where the junction
-		lable	is defined as the CDR3 plus the two flanking conserved
			codons.
junction_aa	string	required, nul-	Amino acid translation of the junction.
		lable	5
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Name	Type .	Attributes	Definition
np1	string	optional, nul-	Nucleotide sequence of the combined N/P region be-
T	6	lable	tween the V gene and first D gene alignment or between
			the V gene and J gene alignments.
np1_aa	string	optional, nul-	Amino acid translation of the np1 field.
1 —		lable	L L
np2	string	optional, nul-	Nucleotide sequence of the combined N/P region be-
1	U U	lable	tween either the first D gene and J gene alignments or
			the first D gene and second D gene alignments.
np2_aa	string	optional, nul-	Amino acid translation of the np2 field.
		lable	
np3	string	optional, nul-	Nucleotide sequence of the combined N/P region be-
Ŧ	U U	lable	tween the second D gene and J gene alignments.
np3_aa	string	optional, nul-	Amino acid translation of the np3 field.
1 —		lable	L L
cdr1	string	optional, nul-	Nucleotide sequence of the aligned CDR1 region.
	6	lable	
cdr1_aa	string	optional, nul-	Amino acid translation of the cdr1 field.
—		lable	
cdr2	string	optional, nul-	Nucleotide sequence of the aligned CDR2 region.
	6	lable	
cdr2_aa	string	optional, nul-	Amino acid translation of the cdr2 field.
—	U U	lable	
cdr3	string	optional, nul-	Nucleotide sequence of the aligned CDR3 region.
		lable	
cdr3_aa	string	optional, nul-	Amino acid translation of the cdr3 field.
		lable	
fwr1	string	optional, nul-	Nucleotide sequence of the aligned FWR1 region.
	_	lable	
fwr1_aa	string	optional, nul-	Amino acid translation of the fwr1 field.
		lable	
fwr2	string	optional, nul-	Nucleotide sequence of the aligned FWR2 region.
		lable	
fwr2_aa	string	optional, nul-	Amino acid translation of the fwr2 field.
		lable	
fwr3	string	optional, nul-	Nucleotide sequence of the aligned FWR3 region.
		lable	
fwr3_aa	string	optional, nul-	Amino acid translation of the fwr3 field.
		lable	
fwr4	string	optional, nul-	Nucleotide sequence of the aligned FWR4 region.
		lable	
fwr4_aa	string	optional, nul-	Amino acid translation of the fwr4 field.
		lable	
v_score	number	optional, nul-	Alignment score for the V gene.
		lable	
v_identity	number	optional, nul-	Fractional identity for the V gene alignment.
		lable	
v_support	number	optional, nul-	V gene alignment E-value, p-value, likelihood, proba-
		lable	bility or other similar measure of support for the V gene
			assignment as defined by the alignment tool.
			Continued on payt page

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Name	Туре	Attributes	Definition
v_cigar	string	required, nul-	CIGAR string for the V gene alignment.
		lable	
d_score	number	optional, nul- lable	Alignment score for the first or only D gene alignment.
d_identity	number	optional, nul-	Fractional identity for the first or only D gene align-
1		lable	ment.
d_support	number	optional, nul- lable	D gene alignment E-value, p-value, likelihood, proba- bility or other similar measure of support for the first or only D gene as defined by the alignment tool.
d_cigar	string	required, nul- lable	CIGAR string for the first or only D gene alignment.
d2_score	number	optional, nul- lable	Alignment score for the second D gene alignment.
d2_identity	number	optional, nul- lable	Fractional identity for the second D gene alignment.
d2_support	number	optional, nul- lable	D gene alignment E-value, p-value, likelihood, proba- bility or other similar measure of support for the second
		luoie	D gene as defined by the alignment tool.
d2_cigar	string	optional, nul- lable	CIGAR string for the second D gene alignment.
j_score	number	optional, nul- lable	Alignment score for the J gene alignment.
j_identity	number	optional, nul- lable	Fractional identity for the J gene alignment.
j_support	number	optional, nul- lable	J gene alignment E-value, p-value, likelihood, probabil- ity or other similar measure of support for the J gene assignment as defined by the alignment tool.
j_cigar	string	required, nul- lable	CIGAR string for the J gene alignment.
c_score	number	optional, nul- lable	Alignment score for the C gene alignment.
c_identity	number	optional, nul- lable	Fractional identity for the C gene alignment.
c_support	number	optional, nul- lable	C gene alignment E-value, p-value, likelihood, proba- bility or other similar measure of support for the C gene assignment as defined by the alignment tool.
c_cigar	string	optional, nul- lable	CIGAR string for the C gene alignment.
v_sequence_star	tinteger	optional, nul- lable	Start position of the V gene in the query sequence (1-based closed interval).
v_sequence_end	integer	optional, nul- lable	End position of the V gene in the query sequence (1-based closed interval).
v_germline_star	tinteger	optional, nul- lable	Alignment start position in the V gene reference se- quence (1-based closed interval).
v_germline_end	integer	optional, nul- lable	Alignment end position in the V gene reference se- quence (1-based closed interval).
v_alignment_sta	ninteger	optional, nul- lable	Start position of the V gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).

Table	2 - continued	from previous page	e
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Name	Туре	Attributes	Definition
v_alignment_end		optional, nul- lable	End position of the V gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
d_sequence_star	_	optional, nul- lable	Start position of the first or only D gene in the query sequence. (1-based closed interval).
d_sequence_end	_	optional, nul- lable	End position of the first or only D gene in the query sequence. (1-based closed interval).
d_germline_star	tinteger	optional, nul- lable	Alignment start position in the D gene reference se- quence for the first or only D gene (1-based closed in- terval).
d_germline_end	integer	optional, nul- lable	Alignment end position in the D gene reference se- quence for the first or only D gene (1-based closed in- terval).
d_alignment_sta	ninteger	optional, nul- lable	Start position of the first or only D gene in both the sequence_alignment and germline_alignment fields (1-based closed interval).
d_alignment_end	-	optional, nul- lable	End position of the first or only D gene in both the sequence_alignment and germline_alignment fields (1-based closed interval).
d2_sequence_sta		optional, nul- lable	Start position of the second D gene in the query se- quence (1-based closed interval).
d2_sequence_end		optional, nul- lable	End position of the second D gene in the query sequence (1-based closed interval).
d2_germline_sta	rinteger	optional, nul- lable	Alignment start position in the second D gene reference sequence (1-based closed interval).
d2_germline_end	1 integer	optional, nul- lable	Alignment end position in the second D gene reference sequence (1-based closed interval).
d2_alignment_st		optional, nul- lable	Start position of the second D gene alignment in both the sequence_alignment and germline_alignment fields (1-based closed interval).
d2_alignment_er		optional, nul- lable	End position of the second D gene alignment in both the sequence_alignment and germline_alignment fields (1-based closed interval).
j_sequence_star		optional, nul- lable	Start position of the J gene in the query sequence (1-based closed interval).
j_sequence_end	_	optional, nul- lable	End position of the J gene in the query sequence (1-based closed interval).
j_germline_star		optional, nul- lable	Alignment start position in the J gene reference se- quence (1-based closed interval).
j_germline_end	_	optional, nul- lable	Alignment end position in the J gene reference sequence (1-based closed interval).
j_alignment_sta	ninteger	optional, nul- lable	Start position of the J gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
j_alignment_end	integer	optional, nul- lable	End position of the J gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
cdr1_start	integer	optional, nul- lable	CDR1 start position in the query sequence (1-based closed interval).
			Continued on post page

Table 2 – continued from previous page

Name	Туре	Attributes	from previous page Definition
cdr1_end	integer	optional, nul-	CDR1 end position in the query sequence (1-based
	-	lable	closed interval).
cdr2_start	integer	optional, nul- lable	CDR2 start position in the query sequence (1-based closed interval).
cdr2_end	integer	optional, nul-	CDR2 end position in the query sequence (1-based
	6	lable	closed interval).
cdr3_start	integer	optional, nul-	CDR3 start position in the query sequence (1-based
	8	lable	closed interval).
cdr3_end	integer	optional, nul-	CDR3 end position in the query sequence (1-based
_	U	lable	closed interval).
fwr1_start	integer	optional, nul-	FWR1 start position in the query sequence (1-based
	6	lable	closed interval).
fwr1_end	integer	optional, nul-	FWR1 end position in the query sequence (1-based
1	integer	lable	closed interval).
fwr2_start	integer	optional, nul-	FWR2 start position in the query sequence (1-based
		lable	closed interval).
fwr2_end	integer	optional, nul-	FWR2 end position in the query sequence (1-based
IWIZ_CIIG	integer	lable	closed interval).
fwr3_start	integer	optional, nul-	FWR3 start position in the query sequence (1-based
IWIJ_SCAIC	Integer	lable	closed interval).
fwr3_end	integer	optional, nul-	FWR3 end position in the query sequence (1-based
IWIJ_end	Integer	lable	closed interval).
fwr4_start	integer	optional, nul-	FWR4 start position in the query sequence (1-based
IWI4_SLAIL	Integer	lable	closed interval).
fwr4_end	intagar	optional, nul-	FWR4 end position in the query sequence (1-based
IWI4_end	integer	lable	closed interval).
v_sequence_alig	ratriang	optional, nul-	Aligned portion of query sequence assigned to the V
	C	lable	gene, including any indel corrections or numbering
			spacers.
v_sequence_alig	n streing t_aa	optional, nul-	Amino acid translation of the v_sequence_alignment
- 1	-	lable	field.
d_sequence_alig	rstreingt	optional, nul-	Aligned portion of query sequence assigned to the first
- 1	-	lable	or only D gene, including any indel corrections or num-
			bering spacers.
d_sequence_alig	n streing t_aa	optional, nul-	Amino acid translation of the d_sequence_alignment
	u –	lable	field.
d2_sequence_ali	gaturinagnt	optional, nul-	Aligned portion of query sequence assigned to the sec-
		lable	ond D gene, including any indel corrections or number-
			ing spacers.
d2_sequence_ali	ostiningnt aa	optional, nul-	Amino acid translation of the d2_sequence_alignment
		lable	field.
j_sequence_alig	ratieng	optional, nul-	Aligned portion of query sequence assigned to the J
	0	lable	gene, including any indel corrections or numbering
			spacers.
j_sequence_alig	natreingt. aa	optional, nul-	Amino acid translation of the j_sequence_alignment
)u		lable	field.
c_sequence_alig	nstreingt	optional, nul-	Aligned portion of query sequence assigned to the con-
a		lable	stant region, including any indel corrections or number-
			ing spacers.
			Continued on next page

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Name Type Attributes Definition c_sequence_alignstreengt_aa optional, nul- Amino acid translation of the c_se	
	quence_angninem
lable field.	
v_germline_aligrstreingt optional, nul- Aligned V gene germline sequence s	
lable region as the v_sequence_alignment	U
the same set of corrections and space	
v_germline_aligrstreingt_aa optional, nul- Amino acid translation of the v_ge	ermline_alignment
lable field.	
d_germline_alignetic optional, nul- Aligned D gene germline sequence s	
lable region as the d_sequence_alignment	
the same set of corrections and space	
d_germline_aligrstreingt_aa optional, nul- Amino acid translation of the d_ge	ermline_alignment
lable field.	
d2_germline_aligstringent optional, nul- Aligned D gene germline sequence s	spanning the same
lable region as the d2_sequence_alignmen	t field and includ-
ing the same set of corrections and sp	bacers (if any).
d2_germline_aligstringent_aa optional, nul- Amino acid translation of the d2_ge	
lable field.	-
j_germline_aligrstreingt optional, nul- Aligned J gene germline sequence s	panning the same
lable region as the j_sequence_alignment t	
the same set of corrections and space	
j_germline_aligrstreingt_aa optional, nul- Amino acid translation of the j_ga	
lable field.	- 0
c_germline_aligrstreingt optional, nul- Aligned constant region germline s	equence spanning
lable the same region as the c_sequence_a	
including the same set of correction	
any).	in and spacers (in
c_germline_aligrstreingt_aa optional, nul- Amino acid translation of the c_g	germline aligment
lable field.	jerninie_unginent
junction_length integer optional, nul- Number of nucleotides in the junction	n sequence
lable	a sequence.
junction_aa_lenginteger optional, nul- Number of amino acids in the junction	on sequence
lable	n sequence.
npl_length integer optional, nul- Number of nucleotides between the	V gene and first D
lable gene alignments or between the V gen	
ments.	te and 5 gene angli
np2_length integer optional, nul- Number of nucleotides between eith	er the first D gene
lable and J gene alignments or the first D g	-
gene alignments.	sene and second D
	acond D cono and
np3_length integer optional, nul- lable J gene alignments.	cond D gene and
	5' of the first or
n1_length integer optional, nul- lable only D gene alignment.	5 of the first of
	2' of the first or
n2_length integer optional, nul- Number of untemplated nucleotides	5 of the first or
lable only D gene alignment.	2^{2} of the second D
n3_length integer optional, nul- Number of untemplated nucleotides	5 of the second D
lable gene alignment.	
p3v_length integer optional, nul- Number of palindromic nucleotides	3' of the V gene
lable alignment.	
	5' of the first or
p5d_lengthintegeroptional, nul-Number of palindromic nucleotideslableonly D gene alignment.	5 of the list of

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Name	Туре	Attributes	Definition
p3d_length	integer	optional, nul-	Number of palindromic nucleotides 3' of the first or
		lable	only D gene alignment.
p5d2_length	integer	optional, nul-	Number of palindromic nucleotides 5' of the second D
		lable	gene alignment.
p3d2_length	integer	optional, nul-	Number of palindromic nucleotides 3' of the second D
		lable	gene alignment.
p5j_length	integer	optional, nul-	Number of palindromic nucleotides 5' of the J gene
		lable	alignment.
consensus_count	integer	optional, nul-	Number of reads contributing to the (UMI) consensus
		lable	for this sequence. For example, the sum of the num-
			ber of reads for all UMIs that contribute to the query
			sequence.
duplicate_count	integer	optional, nul-	Copy number or number of duplicate observations for
		lable	the query sequence. For example, the number of UMIs
			sharing an identical sequence or the number of identical
			observations of this sequence absent UMIs.
cell_id	string	optional, iden-	Identifier defining the cell of origin for the query se-
		tifier, nullable	quence.
clone_id	string	optional, iden-	Clonal cluster assignment for the query sequence.
		tifier, nullable	
repertoire_id	string	optional, iden-	Identifier to the associated repertoire in study metadata.
		tifier, nullable	Identifier to the comple processing chiest in the rener
sample_processi	. rsg <u>ring</u> a	optional, iden- tifier, nullable	Identifier to the sample processing object in the reper- toire metadata for this rearrangement. If the repertoire
		uner, nunable	has a single sample then this field may be empty or miss-
			ing. If the repertoire has multiple samples then this field
			may be empty or missing if the sample cannot be differ-
			entiated or the relationship is not maintained by the data
			processing.
data_processing	ströng	optional, iden-	Identifier to the data processing object in the repertoire
		tifier, nullable	metadata for this rearrangement. If this field is empty
		,	than the primary data processing object is assumed.
rearrangement_i	dstring	DEPRECATED	
	-		tical to sequence_id, but will usually be a universally
			unique record locator for database applications.
rearrangement_s	estr <u>i</u> nigd	DEPRECATED	
germline_databa		DEPRECATED	Source of germline V(D)J genes with version number or
			date accessed.
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Alignment Schema (Experimental)

An Alignment is the output from a V(D)J assignment process for a single V, D, J, or C gene for a sequence. It is not necessary that the V(D)J assignment process performs a sequence alignment algorithm, as the schema can support any algorithmic process. Multiple Alignment records are supported and expected for a single sequence with context-dependent fields (score, identity, support, rank) for assessing the quality of assignments that can vary considerably in definition based on the methodology used.

Note, this schema definition is still experimental and should not be considered final.

File Format Specification

The *format specification* describes the file format and details on how to structure this data.

Fields

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Name	Туре	Attributes	Definition
sequence_id	string	required, nul- lable	Unique query sequence identifier within the file. Most often this will be the input sequence header or a sub- string thereof, but may also be a custom identifier de- fined by the tool in cases where query sequences have been combined in some fashion prior to alignment.
segment	string	required, nul- lable	The segment for this alignment. One of V, D, J or C.
rev_comp	boolean	optional, nul- lable	Alignment result is from the reverse complement of the query sequence.
call	string	required, nul- lable	Gene assignment with allele.
score	number	required, nul- lable	Alignment score.
identity	number	optional, nul- lable	Alignment fractional identity.
support	number	optional, nul- lable	Alignment E-value, p-value, likelihood, probability or other similar measure of support for the gene assign- ment as defined by the alignment tool.
cigar	string	required, nul- lable	Alignment CIGAR string.
sequence_start	integer	optional, nul- lable	Start position of the segment in the query sequence (1-based closed interval).
sequence_end	integer	optional, nul- lable	End position of the segment in the query sequence (1-based closed interval).
germline_start	integer	optional, nul- lable	Alignment start position in the reference sequence (1-based closed interval).
germline_end	integer	optional, nul- lable	Alignment end position in the reference sequence (1-based closed interval).
rank	integer	optional, nul- lable	Alignment rank.
rearrangement_i	.dstring	DEPRECATED	Identifier for the Rearrangement object. May be iden- tical to sequence_id, but will usually be a universally unique record locator for database applications.
data_processing	<u>r s</u> trờng	optional, nul- lable	Identifier to the data processing object in the repertoire metadata for this rearrangement. If this field is empty than the primary data processing object is assumed.
germline_databa	ssering	DEPRECATED	Source of germline V(D)J genes with version number or date accessed.

Clone and Lineage Tree Schema (Experimental)

A unique inferred clone object that has been constructed within a single data processing for a single repertoire and a subset of its sequences and/or rearrangements.

A clone object may have one or more inferred lineage trees. Each tree is represented by a Newick string for its edges and a dictionary of node objects.

File Format Specification

The file format has not been specified yet.

Clone Fields

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Name	Туре	Attributes	Definition
clone_id	string	required, nul- lable	Identifier for the clone.
repertoire_id	string	optional, nul-	Identifier to the associated repertoire in study metadata.
data_processin	g_ströng	lable optional, nul-	Identifier of the data processing object in the repertoire
sequences	array of string	lable optional, nul- lable	metadata for this clone. List sequence_id strings that act as keys to the Rear- rangement records for members of the clone.
v_call	string	optional, nul- lable	V gene with allele of the inferred ancestral of the clone. For example, IGHV4-59*01.
d_call	string	optional, nul- lable	D gene with allele of the inferred ancestor of the clone. For example, IGHD3-10*01.
j_call	string	optional, nul- lable	J gene with allele of the inferred ancestor of the clone. For example, IGHJ4*02.
junction	string	optional, nul- lable	Nucleotide sequence for the junction region of the in- ferred ancestor of the clone, where the junction is de- fined as the CDR3 plus the two flanking conserved codons.
junction_aa	string	optional, nul- lable	Amino acid translation of the junction.
junction_lengt	h integer	optional, nul- lable	Number of nucleotides in the junction.
junction_aa_le	nginteger	optional, nul- lable	Number of amino acids in junction_aa.
germline_align	mestating	required, nul- lable	Assembled, aligned, full-length inferred ancestor of the clone spanning the same region as the se- quence_alignment field of nodes (typically the V(D)J region) and including the same set of corrections and spacers (if any).
germline_align	me stuti<u>n</u>g aa	optional, nul- lable	Amino acid translation of germline_alignment.
v_alignment_st	a rinteger	optional, nul- lable	Start position in the V gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
v_alignment_en	d integer	optional, nul- lable	End position in the V gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
d_alignment_st	arinteger	optional, nul- lable	Start position of the D gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
d_alignment_en	d integer	optional, nul- lable	End position of the D gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
j_alignment_st	annteger	optional, nul- lable	Start position of the J gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
j_alignment_en	d integer	optional, nul- lable	End position of the J gene alignment in both the se- quence_alignment and germline_alignment fields (1- based closed interval).
junction_start	integer	optional, nul- lable	Junction region start position in the alignment (1-based closed interval).
junction_end	integer	optional, nul- lable	Junction region end position in the alignment (1-based closed interval).
aequence_count	integer	optional, nul- lable	Number of Rearrangement records (sequences) in cluded in this clone
seed_id	string	optional, nul- lable	sequence_id of the seed sequence. Empty string (or null) if there is no seed sequence.

Tree Fields

Download as TSV

Name	Туре	Attributes	Definition
tree_id	string	required, nul-	Identifier for the tree.
		lable	
clone_id	string	required, nul-	Identifier for the clone.
		lable	
newick	string	required, nul-	Newick string of the tree edges.
		lable	
nodes	object	optional, nul-	Dictionary of nodes in the tree, keyed by sequence_id
		lable	string

Node Fields

Download as TSV

Name	Туре	Attributes	Definition
sequence_id	string	required, nul-	Identifier for this node that matches the identifier in the
		lable	newick string and, where possible, the sequence_id in
			the source repertoire.
sequence_alignm	nestuting	optional, nul-	Nucleotide sequence of the node, aligned to the
		lable	germline_alignment for this clone, including including
			any indel corrections or spacers.
junction	string	optional, nul-	Junction region nucleotide sequence for the node, where
		lable	the junction is defined as the CDR3 plus the two flank-
			ing conserved codons.
junction_aa	string	optional, nul-	Amino acid translation of the junction.
		lable	

Cell Schema (Experimental)

The cell object acts as point of reference for all data that can be related to an individual cell, either by direct observation or inference.

File Format Specification

The file format has not been specified yet.

Cell Fields

Download as TSV

Name	Туре	Attributes	Definition
cell_id	string	required	Identifier defining the cell of origin for the query se-
			quence.
rearrangements	array of string	required, nul-	Array of sequence identifiers defined for the Rearrange-
		lable	ment object
receptors	array of string	optional, nul-	Array of receptor identifiers defined for the Receptor
		lable	object
repertoire_id	string	required, nul-	Identifier to the associated repertoire in study metadata.
		lable	
data_processing	<u>ströng</u>	optional, nul-	Identifier of the data processing object in the repertoire
		lable	metadata for this clone.
expression_stud	lys <u>t</u> ringthod	optional, nul-	keyword describing the methodology used to assess ex-
		lable	pression. This values for this field MUST come from a
			controlled vocabulary
expression_raw_	dstring	optional, nul-	DOI of raw data set containing the current event
		lable	
expression_inde	string	optional, nul-	Index addressing the current event within the raw data
		lable	set.
expression_tabu	larray of object	optional, nul-	Expression definitions for single-cell
		lable	
virtual_pairing	boolean	required, nul-	boolean to indicate if pairing was inferred.
		lable	

2.3.3 AIRR Software WG - Guidance for AIRR Software Tools

Version 1.0

AIRR Software WG - Compliance Checklist for AIRR Software Tools

Version 1.0 (when finalised)

This questionnaire should be read in conjunction with the AIRR Software WG - Guidance for AIRR Software Tools.

To submit your tool for ratification against the standard, please send the completed questionnaire to software@airrc.antibodysociety.org.

Please provide comments in italics in each response box where these would be helpful to facilitate understanding. We kindly ask for a brief explanatory comment if your answer to a question is *no* or *not applicable*.

Name of Tool:

Contact Name/Institution:

Contact email:

Re- quire- ment Ref.	Question	Response
1	Where is the source code published (please provide a link)?	
2	Does the tool support AIRR Data Representations standards?	yes/no
	Please list any other standard data formats that are supported	
3	Does the distribution include example data?	yes/no
	Is the example data in MiAIRR format, where applicable?	yes/no/not applicable
	Does the tool provide automated checks for expected output from example	yes/no
	data?	
4	Does the output of the tool include a summary of the run parameters?	yes/no
5	Is a container build file provided?	yes/no
	Container technology used?	Docker/Singularity/Othe
	Is the container automatically built as new versions are released?	(please specify)
	Does the automated build run the tool against the example data and test the	yes/no
	output?	yes/no
6	Where can users see what level of support is available? (Please provide a link)	
7	Under what software licence is the tool published? (please provide the name	
	of the licence (e.g. GPL, MIT) or a link	

AIRR Software WG - List of Tools Certified as Compliant

The following tools have been certified as compliant with v1.0 of the guidelines:

Software	Version	Support	Reference
SONAR	3	Output	Schramm et al. Front Immunol,
			2016.
ImmuneDB	0.29.10	Input	Rosenfeld et al. Front Immunol,
			2018,
Scirpy	0.4.2	Input	Sturm et al. Bioinformatics,
			2020,

Evaluation Data Sets

The Software WG is working on the development and evaluation of simulated data sets.

Lists of published real-world datasets are maintained in the AIRR Forum Wiki.

Introduction

The Adaptive Immune Receptor Repertoire (AIRR) Community will benefit greatly from cooperation among groups developing software tools and resources for AIRR research. The goal of the AIRR Software Working Group is to promote standards for AIRR software tools and resources in order to enable rigorous and reproducible immune repertoire research at the largest scale possible. As one contribution to this goal, we have established the following standards for software tools. Authors whose tools comply with this standard will, subject to ratification from the AIRR Software WG, be permitted to advertise their tools as being AIRR-compliant.

Requirements

Tools must:

- 1. Be published in source code form, and hosted on a publicly available repository with a clear versioning system.
- 2. Support community-curated standard file formats and strive for modularity and interoperability with other tools. In particular, tools must read and write *AIRR Data Representations* standards corresponding to their tool.
- 3. Include example data (in AIRR standard formats where applicable) and an automated check for expected output from that data, in order to provide a minimal example of functionality allowing users to check that the software is performing as described.
- 4. Provide information about run parameters as part of the output.
- 5. Provide a container build file that can be used to create an image which encapsulates the software tool, its dependencies, and required run environment. This needs to be remotely and automatically built. The build should conclude by running the example data through the tool (see point 3) and confirming that the expected output is obtained. We currently recognize two software solutions, although we will adapt as software evolves:
 - a. A Dockerfile that automatically builds a container image on Docker Hub.
 - b. A Singularity recipe file that automatically builds a container image on Singularity Hub.
- 6. Provide user support, clearly stating which level of support users can expect, and how and from whom to obtain it.

Recommendations

We suggest software tools be published under a license that permits free access, use, modification, and sharing, such as GPL, Apache 2.0, or MIT. However, we understand that this depends on institutional intellectual property restrictions, thus it is a recommendation rather than a requirement.

Explanatory Notes

Open Source Software and Versioned Repositories

Software tools in the AIRR field are evolving rapidly. In the interests of reproducibility and transparency, published work should be based on tools (and versions of tools) that can be obtained easily by other researchers in the future. To that end, AIRR compliant tools must be published in open repositories such as GitHub or Bitbucket, and we encourage publishing users to provide specifics on the version and configuration of tools that have been employed.

Community-Curated File Formats

The AIRR Data Representation Working Group has defined standards for immune receptor repertoire sequencing datasets. Software tool authors are requested to support these standards as much as possible, for applicable data sets. The currently implemented standard covers submission of reads to NCBI repositories (BioProject, BioSample, SRA and Genbank) and annotated immune receptor rearrangements. Tool authors can assist by easing/guiding the process of submission as much as possible.

Example Data and Checks

Because the installation and operation of the tools in this field may be complex, we require example data and details of expected output, so that users can confirm that their installation is functioning as expected. Furthermore, metadata

(for example, germline gene libraries) and other software dependencies should be checked when the tool runs, and informative error messages issued if necessary. A means should be provided to check the expected output automatically.

Dependencies and Containers

Containers encapsulate everything needed to run a piece of software into a single convenient executable that is largely independent of the user's software environment. For the following purposes, providers of AIRR-compliant tools must provide a containerized implementation (based on a published build script as described above) as one download option that users can choose:

- Containers allow users to use and evaluate a tool easily and reproduce results, without the need to resolve dependencies or configure the environment.
- Having these containers be automatically built also provides a self-validated way to understand the fine details of installation from a known starting point.

To ensure that containers are up to date, they must be built automatically when the current release version of the tool is updated. We will use automated builds on Docker Hub and Singularity Hub for this purpose. The corresponding build files document dependencies clearly, and make it easy for the maintainer to keep the container's dependencies up to date in subsequent releases.

An example Docker container is provided on the Software WG Github Repository. This example encapsulates Ig-BLAST, and implements the bioboxes command-line standard.

Support Statements

Tool authors must provide support for the tool. They must state explicitly what level of support is provided, and explain how support can be obtained. We recommend a method such as the issues tracker on Github, that publishes support requests transparently and links resolutions to specific versions or releases. Users are advised to check that the level of support and the frequency of software updates matches their expectations before committing to a tool.

Analysis Workflows

- At the moment, we do not endorse a specific workflow technology standard:
 - Technology is evolving too rapidly for us to commit to a particular workflow.
 - Typically, AIRR analysis tools have many options and modes, which would make it difficult to support a "plug and play" environment without unduly restricting functionality.
- As tools and workflows evolve, we will keep the position under review and may make stronger technology recommendations in the future.
- We strongly encourage authors of tools to provide concrete, documented, examples of workflows that employ their tools, together with sample input and output data.
- Likewise we encourage authors of research publications to provide documented workflows that will enable interested readers to reproduce the results.

Ratification

Authors may submit tools to the AIRR Software WG requesting ratification against the standard. The submitter should provide a completed copy of the *AIRR Software WG - Compliance Checklist for AIRR Software Tools* to evidence reviewable and itemised evidence of compliance with each Requirement listed above.

The Software WG will, where appropriate, issue a Certificate of Compliance, stating the version of the tool reviewed and the version of the Standard with which compliance was ratified. After receiving a Certificate, authors will be entitled to claim compliance with the Standard, and may incorporate any artwork provided by AIRR for that purpose.

The Software WG will maintain and publish a list of compliant software.

If a tool does not achieve ratification, the Software WG will provide an explanation. The Software WG encourages resubmission once issues have been resolved.

Authors must re-submit tools for ratification following major upgrades or substantial modifications. The Software WG may, at its discretion, request resubmission at any time. If a certified tool subsequently fails ratification, or is not re-submitted in response to a request from the Software WG, AIRR compliance may no longer be claimed and the associated artwork may no longer be used.

The Software WG may, at its discretion, issue a new version of this standard at any time. Tools certified against previous version(s) of the standard may continue to claim compliance with those versions and to use the associated artwork. Authors wishing to claim compliance with the new version must submit a new request for certification and may not claim compliance with the new version, or use associated artwork, until and unless certification is obtained.

2.3.4 AIRR Data Commons API V1

The use of high-throughput sequencing for profiling B-cell and T-cell receptors has resulted in a rapid increase in data generation. It is timely, therefore, for the Adaptive Immune Receptor Repertoire (AIRR) community to establish a clear set of community-accepted data and metadata standards; analytical tools; and policies and practices for infrastructure to support data deposit, curation, storage, and use. Such actions are in accordance with international funder and journal policies that promote data deposition and data sharing – at a minimum, data on which scientific publications are based should be made available immediately on publication. Data deposit in publicly accessible databases ensures that published results may be validated. Such deposition also facilitates reuse of data for the generation of new hypotheses and new knowledge.

The AIRR Common Repository Working Group (CRWG) developed a set of recommendations (v0.6.0) that promote the deposit, sharing, and use of AIRR sequence data. These recommendations were refined following community discussions at the AIRR 2016 and 2017 Community Meetings and were approved through a vote by the AIRR Community at the AIRR Community Meeting in December 2017.

Overview

The AIRR Data Commons (ADC) API provides programmatic access to query and download AIRR-seq data. The ADC API uses JSON as its communication format, and standard HTTP methods like GET and POST. The ADC API is read-only and the mechanism of inclusion of AIRR-seq studies into a data repository is left up to the repository.

This documentation explains how to construct and execute API requests and interpret API responses.

API Endpoints

The ADC API is versioned with the version number (v1) as part of the base path for all endpoints. Each ADC API endpoint represents specific functionality as summarized in the following table:

Endpoint	Туре	HTT	PDescription
/v1	Service status		Returns success if API service is running.
/v1/info	Service information	GET	Upon success, returns service information such as name,
			version, etc.
/v1/	Retrieve a reper-	GET	Upon success, returns the Repertoire information in
repertoire/	toire given its		JSON according to the <i>Repertoire schema</i> .
{repertoire_id}	repertoire_id		
/v1/repertoire	Query repertoires		Upon success, returns a list of Repertoires in JSON
			according to the <i>Repertoire schema</i> .
/v1/	Retrieve a rear-	GET	Upon success, returns the Rearrangement informa-
rearrangement/	rangement given its		tion in JSON format according to the Rearrangement
{sequence_id}	sequence_id		schema.
/v1/	Query rearrangements	POST	Upon success, returns a list of Rearrangements in
rearrangement			JSON or AIRR TSV format according to the Rearrange-
			ment schema.

Authentication

The ADC API currently does not define an authentication method. Future versions of the API will provide an authentication method so data repositories can support query and download of controlled-access data.

Search and Retrieval

The AIRR Data Commons API specifies endpoints for searching and retrieving AIRR-seq data sets stored in an AIRRcompliant Data Repository according to the AIRR Data Model. This documentation describes Version 1 of the API. The general format of requests and associated parameters are described below.

The design of the AIRR Data Commons API was greatly inspired by National Cancer Institute's Genomic Data Commons (GDC) API.

Components of a Request

The ADC API has two classes of endpoints. The endpoints that respond to GET requests are simple services that require few or no parameters. While, the endpoints that response to POST requests are the main query services and provide many parameters for specifying the query as well as the data in the API response.

A typical POST query request specifies the following parameters:

- The filters parameter specifies the query.
- The from and size parameters specify the number of results to skip and the maximum number of results to be returned in the response.
- The fields parameter specifies which data elements to be returned in the response. By default all fields (AIRR and non-AIRR) stored in the data repository are returned. This can vary between data repositories based upon how the repository decides to store blank or null fields, so the fields and/or include_fields parameter should be used to guarantee the existence of data elements in the response.
- The include_fields parameter specifies the set of AIRR fields to be included in the response. This parameter can be used in conjunction with the fields parameter, in which case the list of fields is merged. This is a mechanism to ensure that specific, well-defined sets of AIRR data elements are returned without requiring all of those fields to be individually provided in the fields parameter.

The sets that can be requested are summarized in the table below.

include_fields	MiAIRR	AIRR required	AIRR identifiers	other AIRR fields
miairr	Y	some	N	N
airr-core	Y	Y	Y	N
airr-schema	Y	Y	Y	Y

Service Status Example

The following is an example GET request to check that the service API is available for VDJServer's data repository.

```
curl https://vdjserver.org/airr/v1
```

The response should indicate success.

```
{"result":"success"}
```

Service Info Example

{

The following is an example GET request to get information about the service.

curl https://vdjserver.org/airr/v1

The response provides various information.

```
"name": "adc-api-js-mongodb",
"description": "AIRR Data Commons API reference implementation",
"version": "1.0.0",
"airr_schema_version": 1.3,
"max_size": 1000,
"max_query_size": 2097152,
"contact": {
    "name": "AIRR Community",
    "url": "https://github.com/airr-community"
  }
}
```

Query Repertoire Example

The following is an example POST request to the repertoire endpoint of the ADC API. It queries for repertoires of human TCR beta receptors (filters), skips the first 10 results (from), requests 5 results (size), and requests only the repertoire_id field (fields).

curl --data @query1-2_repertoire.json https://vdjserver.org/airr/v1/repertoire

The content of the JSON payload.

```
{
    "filters":{
        "op":"and",
        "content": [
            {
            "op":"=",
            "content": {
                "field":"subject.organism.id",
                "value":"9606"
            }
        },
        }
}
```

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```
{
    "op":"=",
    "content": {
        "field":"sample.pcr_target.pcr_target_locus",
        "value":"TRB"
     }
     }
     /
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```

The response contains two JSON objects, an Info object that provides information about the API response and a Repertoire object that contains the list of Repertoires that met the query search criteria. In this case, the query returns a list of five repertoire identifiers. Note the Info object is based on the info block as specified in the OpenAPI v2.0 specification.

```
"Info":
{
    "title": "AIRR Data Commons API reference implementation",
    "description": "API response for repertoire query",
    "version": 1.3,
    "contact":
    {
        "name": "AIRR Community",
        "url": "https://github.com/airr-community"
    }
},
"Repertoire":
[
    {"repertoire_id": "4357957907784536551-242ac11c-0001-012"},
    {"repertoire_id": "4476756703191896551-242ac11c-0001-012"},
    {"repertoire_id": "6205695788196696551-242ac11c-0001-012"},
    {"repertoire_id": "6393557657723736551-242ac11c-0001-012"},
    {"repertoire_id": "7158276584776536551-242ac11c-0001-012"}
]
```

Endpoints

The ADC API V1 provides two primary endpoints for querying and retrieving AIRR-seq data. The repertoire endpoint allows querying upon any field in the *Repertoire schema* including study, subject, sample, cell processing, nucleic acid processing, sequencing run, raw sequencing files, and data processing information. Queries on the content of raw sequencing files is not support but is supported on file attributes such as name, type and read information. Queries on Rearrangements is provided by the rearrangement endpoint.

The standard workflow to retrieve all of the data for an AIRR-seq study involves performing a query on the repertoire endpoint to retrieve the repertoires in the study, and one or more queries on the rearrangement endpoint to download the rearrangement data for each repertoire. The endpoints are designed so the API response can be saved directly into a file and be used by AIRR analysis tools, including the AIRR python and R reference libraries, without requiring modifications or transformation of the data.

Repertoire Endpoint

The repertoire endpoint provides access to all fields in the *Repertoire schema*. There are two type of endpoints; one for retrieving a single repertoire given its identifier, and another for performing a query across all repertoires in the data repository.

It is expected that the number of repertoires in a data repository will never become so large such that queries become computationally expensive. A data repository might have thousands of repertoires across hundreds of studies, yet such numbers are easily handled by modern databases. Based upon this, the ADC API does not place limits on the repertoire endpoint for the fields that can be queried, the operators that can be used, or the number of results that can be returned.

Retrieve a Single Repertoire

{

Given a repertoire_id, a single Repertoire object will be returned.

```
curl https://vdjserver.org/airr/v1/repertoire/4357957907784536551-242ac11c-0001-012
```

The response will provide the Repertoire data in JSON format.

```
"Info":
{
   "title": "AIRR Data Commons API reference implementation",
    "description": "API response for repertoire query",
   "version": 1.3,
    "contact":
    {
        "name": "AIRR Community",
        "url": "https://github.com/airr-community"
},
"Repertoire":
[
 {
    "repertoire_id":"4357957907784536551-242ac11c-0001-012",
    "study":{
       "study_id":"PRJNA300878",
       "submitted_by":"Florian Rubelt",
       "pub_ids":"PMID:27005435",
       "lab_name":"Mark M. Davis",
       "lab_address":"Stanford University",
       "study_title":"Homo sapiens B and T cell repertoire - MZ twins"
   },
    "subject":{
       "subject_id":"TW02A",
       "synthetic":false,
       "linked_subjects":"TW02B",
       "organism":{"id":"9606","value":"Homo sapiens"},
       "age":"25yr",
       "link_type":"twin",
       "sex":"F"
   },
    "sample":[
      {"sample_id":"TW02A_T_memory_CD4",
       "pcr_target":[{"pcr_target_locus":"TRB"}],
       "cell_isolation":"FACS",
       "read_length":"300",
       "cell_phenotype":"expression of CD45RO and CCR7",
```

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```
"cell_subset":"Memory CD4+ T cell",
         "filename": "SRR2905669_R1.fastq.gz",
         "single_cell":false,
         "file_type":"fastq",
         "tissue":"PBMC",
         "template_class":"RNA",
         "paired_filename":"SRR2905669_R2.fastq.gz",
         "paired_read_direction":"reverse",
         "read_direction":"forward",
         "sequencing_platform":"Illumina MiSeq"}
      ],
      "data_processing":[
        {"data_processing_id":"4976322832749171176-242ac11c-0001-012",
         "analysis_provenance_id":"651223970338378216-242ac11b-0001-007"}
     1
   }
 ]
}
```

Query against all Repertoires

A query in JSON format is passed in a POST request. This example queries for repertoires of human IG heavy chain receptors for all studies in the data repository.

curl --data @query2_repertoire.json https://vdjserver.org/airr/v1/repertoire

The content of the JSON payload.

```
{
    "filters":{
        "op":"and",
        "content": [
            {
                 "op":"=",
                 "content": {
                     "field":"subject.organism.id",
                     "value":"9606"
                 }
             },
             {
                 "op":"=",
                 "content": {
                     "field":"sample.pcr_target.pcr_target_locus",
                     "value":"IGH"
                 }
            }
        ]
    }
}
```

The response will provide a list of Repertoires in JSON format. The example output is not provided here due to its size.

Rearrangement Endpoint

The rearrangement endpoint provides access to all fields in the *Rearrangement schema*. There are two type of endpoints; one for retrieving a single rearrangement given its identifier, and another for performing a query across all rearrangements in the data repository.

Unlike repertoire data, data repositories are expected to store millions or billions of rearrangement records, where performing "simple" queries can quickly become computationally expensive. Data repositories will need to optimize their databases for performance. Therefore, the ADC API does not require that all fields be queryable and only a limited set of query capabilities must be supported. The queryable fields are described in the Fields section below.

Retrieve a Single Rearrangement

Given a sequence_id, a single Rearrangement object will be returned.

curl https://vdjserver.org/airr/v1/rearrangement/5d6fba725dca5569326aa104

The response will provide the Rearrangement data in JSON format.

```
{
  "Info":
  {
      "title": "AIRR Data Commons API reference implementation",
      "description": "API response for rearrangement query",
      "version": 1.3,
      "contact":
      {
          "name": "AIRR Community",
          "url": "https://github.com/airr-community"
      }
  },
  "Rearrangement":
  [
    {
      "sequence_id":"5d6fba725dca5569326aa104",
      "repertoire_id":"1841923116114776551-242ac11c-0001-012",
      "... remaining fields": "snipped for space"
   }
 ]
}
```

Query against all Rearrangements

Supplying a repertoire_id, when it is known, should greatly speed up the query as it can significantly reduce the amount of data to be searched, though it isn't necessary.

This example queries for rearrangements with a specific junction amino acid sequence among a set of repertoires. A limited set of fields is requested to be returned. The resultant data can be requested in JSON or *AIRR TSV* format.

curl --data @query1_rearrangement.json https://vdjserver.org/airr/v1/rearrangement

The content of the JSON payload.

```
{
    "filters":{
        "op":"and",
        "content": [
            {
            "op":"in",
            "content": {
                "field":"repertoire_id",
                "value":[
                 "2366080924918616551-242ac11c-0001-012",
                "2541616238306136551-242ac11c-0001-012",
                "2541616238306136551-242ac11c-0001-012",
               "2541616238306136551-242ac11c-0001-012",
               "2541616238306136551-242ac11c-0001-012",
               "2541616238306136551-242ac11c-0001-012",
               "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
             "2541616238306136551-242ac11c-0001-012",
              "2541616238306136551-242ac11c-0001-012",
             "2541616238306136551-242ac11c-0001-012",
             "2541616238306136551-242ac11c-0001-012",
             "2541616238306136551-242ac11c-00001-012",
```

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```
"1993707260355416551-242ac11c-0001-012",
                         "1841923116114776551-242ac11c-0001-012"
                    1
                }
            },
            {
                "op":"=",
                "content": {
                    "field":"junction_aa",
                    "value":"CARDPRSYHAFDIW"
                }
            }
       ]
   },
   "fields":["repertoire_id","sequence_id","v_call","productive"],
   "format":"tsv"
}
```

Here is the response in AIRR TSV format.

productive v_call	sequence_id repertoire_id	
true IGHV1-69*04	5d6fba725dca5569326aa106	1841923116114776551-242ac11c-
↔0001-012		
true IGHV1-69*04	5d6fba725dca5569326aa11b	1841923116114776551-242ac11c-
↔0001-012		
true IGHV1-69*10 ↔0001-012	5d6fba725dca5569326aa149	1841923116114776551-242ac11c-
↔0001-012 true IGHV1-69*04	5d6fba735dca5569326aa245	1841923116114776551-242ac11c-
$\Rightarrow 0.001 - 0.012$	Juoiba/JJucaJJ0/J208824J	1041923110114770331 242a0110
true IGHV1-69*04	5d6fba735dca5569326aa274	1841923116114776551-242ac11c-
↔0001-012		
true IGHV1-69*04	5d6fba735dca5569326aa27b	1841923116114776551-242ac11c-
↔0001-012		
true IGHV1-69*04	5d6fba735dca5569326aa27c	1841923116114776551-242ac11c-
↔0001-012		
true IGHV1-24*01	5d6fba735dca5569326aa2a0	1841923116114776551-242ac11c-
↔0001-012 true TGHV1-69*04	5d6fba745dca5569326aa359	1841923116114776551-242ac11c-
rue IGHVI-69*04 $\rightarrow 0001-012$	506102/4500255693262259	1841923116114776551-242aC11C-
true IGHV1-69*04	5d6fba745dca5569326aa408	1841923116114776551-242ac11c-
↔0001-012	545154, 15464555552644400	1011920110111, 00001 242d0110

Request Parameters

The ADC API supports the follow query parameters. These are only applicable to the repertoire and rearrangement query endpoints, i.e. the HTTP POST endpoints.

Parameter	Default	Description
filters	null	Specifies logical expression for query critieria
format	JSON	Specifies the API response format: JSON, AIRR TSV
include_fields	null	Specifies the set of AIRR fields to be included in the response
fields	null	Specifies which fields to include in the response
from	0	Specifies the first record to return from a set of search results
size	repository dependent	Specifies the number of results to return
facets	null	Provide aggregate count information for the specified fields

Filters Query Parameter

The filters parameter enables passing complex query criteria to the ADC API. The parameter represents the query in a JSON object.

A filters query consists of an operator (or a nested set of operators) with a set of field and value operands. The query criteria as represented in a JSON object can be considered an expression tree data structure where internal nodes are operators and child nodes are operands. The expression tree can be of any depth, and recursive algorithms are typically used for tree traversal.

The following operators are support by the ADC API.

Op-	Operan	d\$/alue	Description	Example	
er- a- tor		Data Types			
=	field and value	string, number, inte- ger, or boolean	equals	{"op":"=","content":{"field":"junction_aa","value":"CASSYIKLN"	'}}
!=	field and value	string, number, inte- ger, or boolean	does not equal	{"op":"!=","content":{"field":"subject.organism.id","value":"9606"	}}
<	field and value	number, integer	less than	{"op":"<","content":{"field":"sample.cell_number","value":1000}	
<=	field and value	number, integer	less than or equal	{"op":"<=","content":{"field":"sample.cell_number","value":1000}	}
>	field and value	number, integer	greater than	{"op":">","content":{"field":"sample.cells_per_reaction","value":1(
>=	field and value	number, integer	greater than or equal	{"op":">=","content":{"field":"sample.cells_per_reaction","value":	10000}}
is miss- ing	field	n/a	field is missing or is null	{"op":"is missing","content":{"field":"sample.tissue"}}	
is is	field	n/a n/a	identical to "is missing" opera- tor, provided for GDC compatibil- ity field is not miss-	<pre>{"op":"is","content":{"field":"sample.tissue"}} {"op":"is not missing","content":{"field":"sample.tissue"}}</pre>	
not miss- ing			ing and is not null		
not	field	n/a	identical to "is not missing" op- erator, provided for GDC compat- ibility	{"op":"not","content":{"field":"sample.tissue"}}	
in	field, mul- tiple values in a list	array of string, num- ber, or integer	matches a string or number in a list	{"op":"in","content":{"field":"subject.strain_name","value":["C57B	BL/6","BALB/c
	field, mul- tiple values in a list	array of string, num- ber, or integer	does not match any string or number in a list	{"op":"exclude","content":{"field":"subject.strain_name","value":['SCID","NOD'
-3-A	IRR Star	ndards	contains the sub-	{"op"."contains","content":{"field"."study.study_title","value".99an	cer"}}
tains and	value mul-	n/a	string logical AND	{"op":"and","content":[{"op":"!=","content":{"field":"subject.orga	nism id" "value
unu	tiple	11/ U		{"op":">=","content":{"field":"sample.cells_per_reaction","value":	

Note that the not operator is different from a logical NOT operator, and the logical NOT is not needed as the other operators provide negation.

The field operand specifies a fully qualified property name in the AIRR Data Model. Fully qualified AIRR properties are either a JSON/YAML base type (string, number, integer, or boolean) or an array of one of these base types (some AIRR fields are arrays e.g. study.keywords_study). The Fields section below describes the available queryable fields.

The value operand specifies one or more values when evaluating the operator for the field operand.

Queries Against Arrays

A number of fields in the AIRR Data Model are arrays, such as study.keywords_study which is an array of strings or subject.diagnosis which is an array of Diagnosis objects. A query operator on an array field will apply that operator to each entry in the array to decide if the query filter is satisfied. The behavior is different for various operators. For operators such as = and in, the filter behaves like the Boolean OR over the array entries, that is if **any** array entry evaluates to true then the query filter is satisfied. For operators such as != and exclude, the filter behaves like the Boolean AND over the array entries, that is **all** array entries must evaluate to true for the query filter to be satisfied.

Examples

A simple query with a single operator looks like this:

```
"filters": {
    "op":"=",
    "content": {
        "field":"junction_aa",
        "value":"CASSYIKLN"
    }
}
```

A more complex query with multiple operators looks like this:

```
{
 "filters": {
    "op":"and",
    "content": [
      {
        "op":"!=",
        "content": {
          "field": "subject.organism.id",
          "value":"9606"
        }
      },
      {
        "op":">=",
        "content": {
          "field": "sample.cells_per_reaction",
          "value":"10000"
        }
      },
      {
        "op":"exclude",
        "content": {
          "field": "subject.organism.id",
          "value": ["9606", "10090"]
```

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} } }

Format Query Parameter

Specifies the format of the API response. json is the default format and is available for all endpoints. The rearrangement POST endpoint also accepts tsv which will provide the data in the *AIRR TSV* format. A specific ordering of fields in the TSV format should not be assumed from one API request to another. Take care to properly merge AIRR TSV data from multiple API requests, e.g. such as with the airr-tools merge program.

Fields Query Parameter

The fields parameter specifies which fields are to be included in the API response. By default all fields (AIRR and non-AIRR) stored in the data repository are returned. However, this can vary between data repositories based upon how the repository decides to store blank or null fields, so the fields and/or include_fields parameter should be used to guarantee the existence of data elements in the response.

Include Fields Query Parameter

The include_fields parameter specifies that the API response should include a well-defined set of AIRR Standard fields. These sets include:

- miairr, for only the MiAIRR fields.
- airr-core, for the AIRR required and identifier fields. This is expected to be the most common option as it provides all MiAIRR fields, additional required fields useful for analysis, and all identifier fields for linking objects in the AIRR Data Model.
- airr-schema, for all AIRR fields in the AIRR Schema.

The include_fields parameter is a mechanism to ensure that specific AIRR data elements are returned without requiring those fields to be individually provided with the fields parameter. Any data elements that lack a value will be assigned null in the response. Any empty array of objects, for example subject.diagnosis, will be populated with a single object with all of the object's properties given a null value. Any empty array of primitive data types, like string or number, will be assigned null. Note that if both the include_fields and the fields parameter are provided, the API response will include the set of AIRR fields and in addition will include any additional fields that are specified in the fields parameter.

Size and From Query Parameters

The ADC API provides a pagination feature that limits the number of results returned by the API.

The from query parameter specifies which record to start from when returning results. This allows records to be skipped. The default value is 0 indicating that the first record in the set of results will be returned.

The size query parameters specifies the maximum number of results to return. The default value is specific to the data repository, and a maximum value may be imposed by the data repository. This is to prevent queries from "accidently" returning millions of records. The info endpoint provides the data repository default and maximum values for the repertoire and rearrangement endpoints, which may have different values. A value of 0 indicates there is no limit on the number of results to return, but if the data repository does not support this then the default value will be used.

The combination of from and size can be used to implement pagination in a graphical user interface, or to split a very large download into smaller batches. For example, if an interface displays 10 records as a time, the request would assign size=10 and from=0 to get the ten results to display on the first page. When the user traverses to the "next page", the request would assign from=10 to skip the first ten results and return the next ten results, and from=20 for the next page after that, and so on.

Facets Query Parameter

The facets parameter provides aggregate count information for the specified field. Only a single field can be specified. The facets parameter can be used in conjunction with the filters parameter to get aggregate counts for a set of search results. It returns the set of values for the field, and the number of records (repertoires or rearrangement) that have this value. For field values that have no counts, the API service can either return the field value with a 0 count or exclude the field value in the aggregation. The typical use of this parameter is for displaying aggregate information in a graphical user interface.

Here is a simple query with only the facets parameter to return the set of values for sample.pcr_target. pcr_target_locus and the count of repertoires repertoires that have each value. The content of the JSON payload.

```
"facets":"sample.pcr_target.pcr_target_locus"
```

Sending this query in an API request.

}

```
curl --data @facets1_repertoire.json https://vdjserver.org/airr/v1/repertoire
```

The output from the request is similar to normal queries except the data is provided with the Facet key.

```
{
  "Info": {
    "title": "AIRR Data Commons API reference implementation",
    "description": "API response for repertoire query",
    "version": 1.3,
    "contact": {
        "name": "AIRR Community",
        "url": "https://github.com/airr-community"
     }
    },
    "Facet": [
        {"sample.pcr_target.pcr_target_locus":[["TRB"]],"count":40},
        {"sample.pcr_target.pcr_target_locus":[["IGH"]],"count":20}
    ]
}
```

Here is a query with both filters and facets parameters, which restricts the data records used for the facets count. The content of the JSON payload.

```
{
    "filters":{
        "op":"=",
        "content": {
            "field":"sample.pcr_target.pcr_target_locus",
            "value":"IGH"
        }
    },
    "facets":"subject.subject_id"
}
```

Sending this query in an API request.

curl --data @facets2_repertoire.json https://vdjserver.org/airr/v1/repertoire

Example output from the request. This result indicates there are ten subjects each with two IGH repertoires.

```
"Info": {
   "title": "AIRR Data Commons API reference implementation",
   "description": "API response for repertoire query",
   "version": 1.3,
   "contact": {
     "name": "AIRR Community",
      "url": "https://github.com/airr-community"
   }
 },
  "Facet": [
   {"subject.subject_id":"TW05B","count":2},
   {"subject.subject_id":"TW05A","count":2},
   {"subject.subject_id":"TW03A","count":2},
    {"subject.subject_id":"TW04A","count":2},
    {"subject.subject_id":"TW01A","count":2},
    {"subject.subject_id":"TW04B","count":2},
   {"subject.subject_id":"TW02A","count":2},
   {"subject.subject_id":"TW03B","count":2},
   {"subject.subject_id":"TW01B","count":2},
    {"subject.subject_id":"TW02B","count":2}
 ]
}
```

ADC API Limits and Thresholds

Repertoire endpoint query fields

It is expected that the number of repertoires in a data repository will never become so large such that queries become computationally expensive. A data repository might have thousands of repertoires across hundreds of studies, yet such numbers are easily handled by databases. Based upon this, the ADC API does not place limits on the repertoire endpoint for the fields that can be queried or the operators that can be used.

Rearrangement endpoint query fields

Unlike repertoire data, data repositories are expected to store billions of rearrangement records, where performing "simple" queries can quickly become computationally expensive. Data repositories are encouraged to optimize their databases for performance. Therefore, based upon a set of query use cases provided by immunology experts, a minimal set of required fields was defined that can be queried. These required fields are described in the following Table. The fields also have the AIRR extension property adc-query-support: true in the AIRR Schema.

Field(s)	Description		
sequence_id, reper-	Identifiers; sequence_id allows for query of that specific rearrangement object in the		
toire_id, sam-	repository, while repertoire_id, sample_processing_id, and data_processing_id are links		
ple_processing_id,	to the repertoire metadata for the rearrangement. The clone_id and cell_id are identifiers		
data_processing_id,	that group rearrangements based on clone assignment and single cell assignment.		
clone_id, cell_id			
locus, v_call, d_call,	Commonly used rearrangement annotations.		
j_call, c_call, produc-			
tive, junction_aa, junc-			
tion_aa_length			

Data repository specific limits

A data repository may impose limits on the size of the data returned. This might be because of limitations imposed by the back-end database being used or because of the need to manage the load placed on the server. For example, MongoDB databases have document size limits (16 megabytes) which limit the size of a query that can be sent to a repository and the size of a single repertoire or rearrangement object that is returned. As a result a repository might choose to set a maximum query size.

Size limits can be retrieved from the info endpoint. If the data repository does not provide a limit, then no limit is assumed.

Field	Description		
max_siz	eThe maximum value for the size query parameter. Attempting to retrieve data beyond this maximum		
	should trigger an error response. The error response should include information about why the query		
	failed and what the maximum size limit is.		
max_que	max_querFhesmaximum size of the JSON query object.		

Reference Implementation

The AIRR Community provides a reference implementation for an ADC API service with more information found *here*.

2.3.5 AIRR Ontologies and Vocabularies Team

Summary

The "Ontologies and Vocabularies Team" was formed as a joint interest group of the Common Repository (ComRepo) and the Minimal Standards (MiniStd) working groups of the AIRR Community. The long-term aim of the Team is to define standard vocabularies and ontologies to be used by AIRR-compliant databases.

Ontology Data Representation

The nodes in an ontology are typically either concepts (e.g., capital) or instances thereof (e.g., Paris). These nodes have *local IDs* (often numbers), which are unique within an ontology. They also typically have *labels*, which is the human-readable name of the node. Ontology entities in the AIRR Data Standard reflect this model, with each AIRR field that is represented as an ontology recorded with a global *ontology ID* (id) and the corresponding *label* (label).

Within the AIRR Standards, Compact URIs (CURIEs) are used to represent *ontology IDs*. CURIEs are a standardized way to abbreviate International Resource Identifiers (IRI, [RFC3987]), which includes URIs as a subset. They were originally conceived to simplify the handling of attributes, e.g. in XML or SPARQL, by making them more compact and readable. CURIEs are also used by IEDB databases to reduce redundancies (mainly in the leading part of IRIs).

For example, a typical CURIE would look like NCBITAXON: 9258. In this case, NCBITAXON is the *prefix*, a custom string that will be replaced by a repository-defined IRI component (e.g., http://purl.obolibrary.org/obo/NCBITaxon_). Note that there is no connection between NCBITAXON in the CURIE and NCBITAXON in the IRI, the former one is just a placeholder.

The AIRR schema will provide a list of AIRR approved CURIE *prefixes* along with a list of at least one IRI *prefix* (i.e., replacement string) for each them. This list serves two purposes:

- 1. It provides a controlled namespace for CURIE *prefixes* used in the AIRR schema. For now, custom additions to or replacements of these *prefixes* in the schema are prohibited. This does not affect the ability of repositories to use such custom prefixes internally.
- 2. It simplifies resolution of CURIEs by non-repositories. The lists of IRI *prefixes* for each CURIE *prefix* should not be considered to be exhaustive. However, when using custom IRI *prefixes*, it must be ensured that they refer to the same ontology as the provider *prefixes*.

It should be explicitly noted that the IRI *prefix* list should not be interpreted as any kind of recommendation for certain *providers*. It is left up to users to decide how to resolve the resulting IRIs, e.g., via DNS/HTTP (if possible) or by using a *provider* of their choice.

Approved Ontologies

- Cell ontology (CL)
 - used in:
 - * Cell subset (cell_subset, Tissue and Cell Processing)
 - CURIE summary
 - * CURIE Prefix: CL
 - * CURIE IRI Prefix: http://purl.obolibrary.org/obo/CL_
 - example AIRR use
 - * "cell_subset.id" : "CL:0000542"
 - * "cell_subset.label" : "lymphocyte"
 - default root node
 - * label: lymphocyte
 - * local id: CL_0000542
 - * path: "
 - license: CC BY
 - latest release (as of 2020-05-20): 2020-03-02
 - repo: https://github.com/obophenotype/cell-ontology
 - maintainer: Alexander Diehl, Buffalo, NY, US (addiehl@buffalo.edu)
- Human disease ontology (DOID)
 - used in:
 - * Diagnosis (disease_diagnosis, Diagnosis)
 - CURIE summary
 - * CURIE Prefix: DOID
 - * CURIE IRI Prefix: http://purl.obolibrary.org/obo/DOID_
 - example AIRR use
 - * "disease_diagnosis.id" : "DOID:9538"
 - * "disease_diagnosis.label" : "multiple myeloma"
 - default root node
 - * label: disease
 - * local ID: DOID: 4
 - * path: disease
 - license: CC0
 - latest release (as of 2020-05-20): 2020-04-20

- repo: https://github.com/DiseaseOntology/HumanDiseaseOntology
- maintainer: Lynn Schriml, U Maryland, MD, US (lynn.schriml@gmail.com)
- notes: Features ICD cross-reference
- NCBI organismal taxonomy (NCBITAXON)
 - used in:
 - * Species (species, Subject)
 - * Cell species (cell_species, Tissue and Cell Processing)
 - CURIE summary
 - * CURIE Prefix: NCBITAXON
 - * CURIE IRI Prefixes: http://purl.obolibrary.org/obo/NCBITaxon_, http:// purl.bioontology.org/ontology/NCBITAXON/
 - example AIRR use
 - * "species.id": "NCBITAXON:9606"
 - * "species.label" : "Homo sapiens"
 - default root node
 - * label: Gnathostomata
 - * local ID: 7776
 - * path: cellular organisms/Eukaryota/Opisthokonta/Metazoa/Eumetazoa/ Bilateria/Deuterostomia/Chordata/Craniata/Vertebrata/Gnathostomata
 - license: UMLS
 - latest release (as of 2020-05-20): 2020-04-18
 - repo: https://github.com/obophenotype/ncbitaxon
 - maintainer: NCBI (info@ncbi.nlm.nih.gov)
- NCI thesaurus (NCIT)
 - used in:
 - * Study type (study_type, Study)
 - CURIE summary
 - * CURIE Prefix: NCIT
 - * CURIE IRI Prefixes: http://purl.obolibrary.org/obo/NCIT_, http://ncicb. nci.nih.gov/xml/owl/EVS/Thesaurus.owl#
 - example AIRR use
 - * "study_type.id" : "NCIT:C15197"
 - * "study_type.label" : "Case-Control Study"
 - default root node
 - * label: Study
 - * local ID: C63536
 - * path: Activity/Clinical or Research Activity/ Research Activity/Study

- license: Public domain, credit of NCI is requested
- repo: https://github.com/NCI-Thesaurus/thesaurus-obo-edition
- latest release (as of 2020-05-20): 2020-05-04
- maintainer: NCI (ncicbiitappssupport@mail.nih.gov)
- Units of measurement ontology (UO)
 - used in:
 - * Age unit (age_unit, Subject)
 - CURIE summary
 - * CURIE Prefix: UO
 - * CURIE IRI Prefix: http://purl.obolibrary.org/obo/U0_
 - example AIRR use
 - * "age_unit.id" : "UO:0000036"
 - * "age_unit.label" : "year"
 - default root node
 - * label: time unit
 - * local ID: UO_000003
 - * path: unit/time unit
 - license: CC BY (per Github repo)
 - repo: https://github.com/bio-ontology-research-group/unit-ontology
 - latest release (as of 2020-05-20): 2020-05-18
 - maintainer: unknown
- Uber-anatomy ontology (Uberon)
 - used in:
 - * Tissue (tissue, Sample)
 - CURIE summary
 - * CURIE Prefix: UBERON
 - * CURIE IRI Prefix: http://purl.obolibrary.org/obo/UBERON_
 - example AIRR use
 - * "tissue.id" : "UBERON:0002371"
 - * "tissue.label" : "bone marrow"
 - default root node
 - * label: multicellular anatomical structure
 - * local ID: UBERON:0010000
 - * path: /BFO_0000002/BFO_0000004/anatomical entity/material anatomical entity/anatomical structure/multicellular anatomical structure
 - license: CC BY

- repo: https://github.com/obophenotype/uberon
- latest release (as of 2020-05-20): 2019-11-22
- maintainer: Chris Mungall, LBL, CA, US (cjmungall@lbl.gov)

Sprint Reports

OntoVoc Report - Sprint 11/2018

Objectives

The objectives of this first sprint in November 2018 were to:

- 1. define criteria for suitable ontologies
- 2. identify ontologies for five fields/keywords of the MiAIRR data standard and
- 3. assess technical aspects of ontology integration into databases

General Considerations

The Team initially discussed an approach where only vocabularies (i.e. lists of terms) and not ontologies (i.e. many terms connected by predicates) would have been defined. These vocabularies would have been derived from ontologies, but this process would not necessarily have been reversible. The notion at this time point was, that such an approach would allow to solve a number of problems like combining multiple sources and removing duplicated leaves. However, after some discussions this approach was effectively abandoned for a number of reasons:

- It would discard the UID for an entity. As the UID (in contrast to the name string) is guaranteed to be stable and unique, it facilitates updates, linking and information representation, all of which would otherwise be lost.
- In general, it will be more sustainable to work with the maintainers of an existing ontology to include entities/terms, than just dumping their terms into a list and adding new ones.
- Well-designed ontologies will not contain duplicated entities, although they might appear to do so in a simple browsers (i.e. this is an artifact of representation). Ontologies that actually do contain duplicates are excluded by *criterium 2*.

Criteria for Ontologies

Criteria

Ontologies used within AIRR standards

- 1. MUST¹ cover the majority of the required terms, but complete coverage is OPTIONAL
- 2. MUST have a structure that is scientifically correct and logically coherent
- 3. MUST NOT feature complexity that makes it hard to use for queries and data representation
- 4. SHOULD already be widely adopted
- 5. MUST be actively maintained
- 6. MUST be available under a free license

¹ See the "Glossary" section on how to interpret term written in all-caps.

Comments on criteria:

- ad (1): For most fields it will be difficult to find complete and accurate ontologies. Therefore picking the best available ontology and working with its maintainers to include missing terms is expected to be the most sustainable approach.
- ad (5): This requirement follows from (1), as there needs to be a way for term requests.
- ad (6): A number of ontologies need to be licensed from their respective copyright holders. This results in potential barriers for implementation and distribution of such ontologies. Therefore only ontologies available under a free license are considered suitable for AIRR-compliant databases. The list of suitable licenses is not final, but includes: CC0 and CC BY.

Selected Ontologies

(designations are MiAIRR field names and DataRep keywords)

Completed

- Species (organism)
 - NCBITAXON
 - license: UMLS²
 - latest release: 2018-07-06
 - maintainer: NCBI (info@ncbi.nlm.nih.gov)
- Diagnosis (disease_diagnosis)
 - DOID
 - root node
 - * name: disease
 - * ID: DOID:4
 - * path: /disease
 - License: CC BY
 - latest release: 2018-03-02
 - maintainer: Lynn Schriml, U Maryland, MD, US (lynn.schriml@gmail.com)
 - notes: Features ICD cross-reference
- Cell subset (cell_subset)
 - CellOntology
 - license: CC BY
 - latest release: 2018-07-11
 - maintainer: Alexander Diehl, Buffalo, NY, US (addiehl@buffalo.edu)
- Tissue (tissue)

⁻ Uberon

 $^{^{2}}$ Will require further review the UMLS Metathesaurus License is not a free license, however it needs to be clarified how much of it relates to the work (i.e. the taxonomy itself) and how much to the service.

- root node
 - * name: multicellular anatomical structure
 - * **ID**: UBERON: 0010000
 - * path: /BFO_0000002/BFO_0000004/anatomical entity/material anatomical entity/anatomical structure/multicellular anatomical structure
- License: CC BY
- latest release: 2018-10-15
- Maintainer: Chris Mungall, LBL, CA, US (cjmungall@lbl.gov)

Under evaluation

- Strain name (strain_name)
 - Suggested ontologies:
 - * JAX
 - * IEDB
 - Issues:
 - * Nomenclature
 - * one ontology is not enough

Technical aspects

- Repositories:
 - UID assigned by ontologies are guaranteed to be unique and permanent³.
 - A repository MAY use internal identifiers that are distinct from UIDs. However, to be AIRR-compliant it MUST be able to map UIDs to its identifiers.
 - Points of "AIRR compliance" would typically be:
 - * When data is extracted from the repository through a Query API (CRWG)
 - * When data is extracted from the repository into a file format (DataRep)
- Integration of ontologies into repositories:
 - There are two main ontology providers offering a REST API and all the ontologies listed above:
 - * NCBO Bioportal
 - * OLS ontology
 - NCBO can apparently be slow and sometimes not that stable, while OLS seems to be more stable and
 potentially has a better long-term support.
 - Remote ontology services tend to be slow and create external dependencies. On the other hand, while local
 hosting of an ontology is possible (and partially supported by NCBO and OLS), it requires non-negligible
 resources. The Team's current assumption is that queries to remote ontology services can be substantially

 $^{^{3}}$ This has more recently (early 2020) been called in question and will be revisited during the next sprint. Note that the uncertainty revolves around the question what exactly constitues a UID, rather than the question whether a UID is permanent and unique.

accelerated if only the relevant section of a respective ontology is queried. Therefore a local service would not be necessary.

- Repositories should store both the IDs and the values in their database. This way, they do not have to query the ontology in a scenario where human-readable output is required. In the case of changes, most ontologies try to follow the practice of not changing a term value but instead create a new term with the new value and a new ID, and deprecating the old term. Therefore term deprecation needs to be handled by the repository.
- Like for the databases, also the API should be able to handle both IDs and values as query input and return both during a query.
- The user interface (UI) should offer an ontology-backed autocomplete. NCBO provides some JavaScript code to use. The UI must not offer deprecated terms. To allow entry of terms not present in the ontology, data can be prefixed with some text that will allow the data validation to proceed (e.g., if an entry starts with "other -" the UI will not autocomplete/validate). Later, i.e. when the term has been created, the data will be updated.
- Note that the complete IEDB can be downloaded as SQL dump, it is licensed under CC BY. At a first glance, the main overlap seems to be with organism, strain_name and to a smaller extent disease_diagnosis. However, sample information like cell_subset and tissue seems to be largely absent from IEDB, so it could currently not be the one-stop solution for AIRR.

Footnotes

Appendix

Glossary

- MUST / REQUIRED: Indicates that an element or action is necessary to conform to the standard.
- SHOULD / RECOMMENDED: Indicates that an element or action is considered to be best practice by AIRR, but not necessary to conform to the standard.
- MAY / OPTIONAL: Indicates that it is at the discretion of the user to use an element or perform an action.
- MUST NOT / FORBIDDEN: Indicates that an element or action will be in conflict with the standard.

OntoVoc Report - Sprint 04/2020

Objectives

The objectives of this second sprint in April 2020 were to:

- 1. revisit general policies around ontologies used in the AIRR schema
- 2. identify two new ontologies for several fields of the AIRR schema
- 3. solve technical questions regarding IDs and providers

General Policies

The OntoVoc team revisited the criteria for ontologies used in the AIRR schema that it *defined in the 11/2018 sprint*. While they are still considered to be valid, the team felt that a more detailed guidance could be useful in the process of selecting ontologies for new fields. It therefore evaluated the OBO Foundry Principles, which partially re-iterate

some of the existing criteria (e.g., *Openness* and *Maintenance*), but also provide additional recommendations, e.g., the presence of textual definitions, clear scope and a common format, which were considered to be valuable additions to the existing guidelines. The team therefore decided to endorse the OBO Foundry Principles, as RECOMMENDED (but NOT REQUIRED) criteria. It should be noted, that this does not make any statement regarding the use of OBO vs. non-OBO ontologies.

Decisions on Pending Items of Sprint 11/2018

A number of decisions on draft and legacy ontologies as well as root nodes was not officially passed during the last sprint. The team thus revisited and confirmed the following decisions:

- Use of NCIT for study_type, top node Study (NCIT:C63536).
- Use of UO for age_unit, top node time unit (UO:000003).
- Use of `Gnathostomata (NCBITAXON: 7776) as top node for NCBITAXON when used for fields encoding a host species.
- Use of lymphocyte (CL:0000542) as top node for CL when used for cell_subset.

New Ontologies

Mouse strain

Background

Mouse strain names follow a very elaborate nomenclature that is capable of describing the genetic background, breeding history and introduced mutation in a detailed manner. However, this nomenclature is rarely used correctly (if at all), which creates uncertainty about the identity of strains used in experimental studies. Therefore an ontology or vocabulary compliant to this nomenclature would be of tremendous help for consistent annotation.

An ontology for the strain_name field was already on the list for the last sprint, however it was not possible to identify a single ontology that would contain comprehensive information about strains from multiple species. This situation created a problem that could not be resolved then. In the meantime, the concept of "extensions" has been introduced to the AIRR schema, which create an additional layer of fields (and associated ontologies) on top of a core schema. As these extensions can be made conditional on the value of fields within the core schema, it has now become possible to have multiple extensions defining the strain_name field, but for different species and therefore with distinct species-specific ontologies.

Having addressed this issue, the other key problem that remains is the absence of an actual ontology for mouse strains, while a rat strain ontology exists. Therefore in a first step it is necessary to identify resources that you at least serve as a provider for vocabularies. The two potential candidates that were identified are:

- MGI: The Mouse Genome Informatics database hosted at JAX aims to be comprehensive in regard to all mouse strains that have been published in the literature.
- IEDB: The Immune Epitope Database already ran into the problem of a missing mouse ontology and therefore decided to build up their own reference focused on immunologically relevant strains, as part of their Ontie database.

Once it is clear which of the resources could be used, it will be necessary to approach the current maintainers regarding their willingness to convert the data into an actual ontology (the RS could serve as a template for this). As this will take longer than just a couple of weeks, the second step is out-of-scope for this sprint.

Evaluation

- MGI: The database can be downloaded as a dump, however the licensing conditions are unclear. It contains a total of 60k entries of which 3.2k inbred and 13.8k are congenic strains. The majority of the remaining entries are coisogenic strains, most of them from large- scale gene KO projects.
- IEDB: Database dumps can also be downloaded and are freely available under CC-BY 4.0. It covers over a thousand mouse strains and contains additional information on the genetic background of a strain.

Next steps

• Get in touch with JAX (pending)

Geolocation

There are several (planned) extensions to the AIRR metadata standard that will provide geospatial metadata. Countrylevel information is typically assumed to be privacy-preserving and easy to operationalize. Therefore, while clearly only capturing some aspects of genetic ancestry, it might serve as a proxy for concepts of "race" and "ethnicity" that are rather ill-defined.

Potential candidate vocabularies/ontologies:

- ISO3166-1 alpha-2: Two-letter code, some ambiguity but well known from ccTLDs.
- ISO3166-1 alpha-3: Three-letter code, less ambiguity than alpha-2.
- UN Stats Division code (currently M49): Numerical code, not human-readable, maps to ISO3166-1 alpha-3.
- Gazetter (GAZ)
 - Contains 2nd (state) and 3rd (county) level information.
 - Not linked to any actual coordinates
 - ISO3166-1 annotation is incomplete and lacks e.g. for Germany and Switzerland.
 - Does not support German Umlauts. Äbsölütely inacceptable, as these are not just diacritical marks (i.e. "Münster" and "Munster" are two different cities).
- HANCESTRO:
 - Seems to be complete, but does not provide ISO3166 codes.
 - Ontology could also be used for other fields relating to genetic ancestry.
 - Links to DBpedia, currently unclear whether it is also populated from there
 - country node has pan-240 leaves (surplus seems due to oversea territories), cross-referencing to GAZ (s/a)
- Various pathogen-related repositories:
 - VectorBase (VBGEO): see link and choose "GADM/VBGEO PlaceNames"
 - Viral Pathogen Resource (ViPR):
 - * Uses v1.3 of the GSCID/BRC Project and Sample Application Standard.
 - * GSCID/BRC Core Sample defines four fields for "Collection Location":
 - "Latitude" (CS11) and "Longitude" (CS12) in ISO 6709 format
 - $\cdot\,$ "Location" (CS13), using GAZ as controlled vocabulary

• "Country" (CS14) as by ISO3166-1 (alpha-2).

- Influenza Research Database (IRD): Flu-focused version of ViPR, also uses GSCID/BRC Project and Sample Application Standard v1.3.
- Pathosystems Resource Integration Center (Patric): Focused on bacterial infectious diseases. Uses an "Isolation Country" field in their "Genome" table, format seems to be full text.

Rejected candidates:

- HL7: own ontology deprecated, now recommends ISO 3166-1 alpha-3 set.
- NCIT: Incomplete, only contains pan-90 entities
- SNOMED: Licensing issues
- GADM data: Good quality and resolution, but not an ontology in itself. Also not under a free license, does not allow redistribution or commercial use.

Evaluation

Given the number of options, there is no obvious candidate to pick. Therefore the team decided to define clear use cases and then evaluate each options against them. However, due to time limitation, we did not really get into this, will have to follow up in the next sprint. The use cases so far were:

- Annotate country of birth / of sampling [REQUIRED]
- Encode higher resolution than country level if legally permitted and scientifically meaningful [RECOM-MENDED].
- Linking to geo-spatial coordinates [OPTIONAL]

Technical Questions

Background and Problem

Some nomenclature first: The nodes in an ontology graph are typically either *concepts* (e.g., capital) or *instances* thereof (e.g., Paris). These nodes have *local IDs* (often numbers), which are unique within an ontology. They also typically have *labels*, which is the human- readable name of the node. Nodes can have additional *attributes* (e.g., "population count") and are connected to other nodes by *relations* (e.g. "is-a", "superset-of"), which create the edges of the graph.

The complete ontology is usually represented in an XML or OWL file. However, we are looking for a *provider*, i.e. a service that facilitates queries of an ontology via web and/or an API-based interface. Upon querying with a unique ID, is it expected that a *provider* will be able to return the record of a node, which should contain all attributes and relations. Furthermore a *provider* might allow set- and graph-based queries (e.g., is A a complete subset of B; what is the last common ancestor of A and B). Finally a *provider* can offer lookup services, i.e., identify the corresponding *concept* or *instance* in another ontology. Until now we have mainly looked at three providers: Ontobee, OLS and BioPortal. While they all provide similar basic services, it should be noted that some biomedical databases and repositories are, by convention, restricted to use certain *providers*.

As stated above, each node has a *local ID*. To avoid conflicts between the *local IDs* of multiple ontologies, *providers* and ontology collections (e.g., OBO Foundry) use a namespace, i.e., some abbreviation for the ontology that is prefixed to the *local ID*. However, as there no common standard how to create these prefixes, this system is only unambiguous and collision-safe within a single *provider*. To resolve this issue, ontologies often use International Resource Identifiers (IRI, [RFC3987]). While IRIs look like HTTP URLs, they should primarily be considered as permanent and globally unique identifiers, which might resolve to the node's record via DNS/HTTP, but this is optional. In addition, potential intermediate URLs generated in the DNS/HTTP resolving process must be considered internal and therefore should

not be used by third parties. Finally, it needs be noted that IRIs should to be considered case-sensitive, especially when used as identifiers (per [RFC3987], Section 5.3.2.1, which only excludes the schema and host (authority) component for case-sensitivity).

While many ontologies already define an entities IRI on the level of the ontology, there are some that do not. For such ontologies, IRIs are then assigned by the provider. The most notable example for this are the UMLS ontologies like the NCBI Taxonomy. This leads to the situation that a single node in an ontology, stored by two providers can have different IRIs. Therefore, a concept from NCBI Taxonomy, e.g., the duck-billed platypus (label: *Ornithorhynchus anatinus*, local ID: 9258) has the IRI http://purl.obolibrary.org/obo/NCBITaxon_9258 in Ontobee and the IRI http://purl.bioontology.org/ontology/NCBITAXON/9258 in BioPortal. In addition, other providers might choose to use one of these IRIs too, although it will never resolve to their system via DNS/HTTP (e.g., OLS uses the Ontobee IRIs).

For the AIRR Community, this creates the challenge that we want to be able to have unambiguous identifiers, without requiring any specific provider.

Proposed solution

Compact URIs (CURIEs) are a standardized way to abbreviate IRIs, which includes URIs as a subset. They were originally conceived to simplify the handling of attributes, e.g. in XML or SPARQL, by making them more compact and readable. CURIEs are e.g. used by IEDB databases to reduce redundancies (mainly in the leading part of IRIs).

A typical CURIE would, e.g., look like NCBITAXON:9258. In this case, NCBITAXON is the *prefix*, a custom string that will be replaced by a repository-defined IRI component (e.g., http://purl.obolibrary.org/obo/NCBITaxon_). Note that there is no connection between NCBITAXON in the CURIE and NCBITAXON in the IRI, the former one is just a placeholder.

This resolves the issue of different *providers* usings different IRIs with distinct formatting rules (as described above). As the choice of the *provider* is independent for each ontology, it allows greater flexibility for the repositories, as they do not need a single *provider* that needs be able to resolve all terms. Similarly, different repositories can use the same ontology, but use different *providers*. Note that this would not require changes to the data, as the data would only contains CURIEs, not the (provider-specific) IRIs.

The AIRR schema will provide a list of AIRR approved CURIE *prefixes* along with a list of at least one IRI *prefix* (i.e., replacement string) for each them. This list serves two purposes:

- 1. It provides a controlled namespace for CURIE *prefixes* used in the AIRR schema. For now, custom additions to or replacements of these *prefixes* in the schema are prohibited. This does not affect the ability of repositories to use such custom prefixes internally.
- 2. It simplifies resolution of CURIEs by non-repositories. The lists of IRI *prefixes* for each CURIE *prefix* should not be considered to be exhaustive. However, when using custom IRI *prefixes*, it must be ensured that they refer to the same ontology as the provider *prefixes*.

It should be explicitly noted that the IRI *prefix* list should not be interpreted as any kind of recommendation for certain *providers*. It is left up to users to decide how to resolve the resulting IRIs, e.g., via DNS/HTTP (if possible) or by using a *provider* of their choice.

Modifications to the AIRR schema

All changes to the AIRR schema that would be based on the sprint can currently be reviewed on Github in Pull Request #385. These changes are intended to be included into the next major release.

Clarifications

- Root nodes are specific to individual fields, not to an ontology. Therefore, NCBITAXON will use a root node of "Gnathostomata" for the annotations of the host species, but this would not be useful, e.g., if it would be used to annotate pathogenic organisms, which will require a top node at the apex of the hierarchy.
- The labels (previous: values) that are provided in the schema for ontology-based fields, should be considered an addition for convenience and not as being authoritative. Repositories or applications can choose to link synonyms to given concepts (e.g., "human" for "*Homo sapiens*") to simply search queries. Repositories further can provide such a synonym in the label field upon exporting data. However, repositories importing data should verify the correctness of labels that do not match the one provided by the ontology. Importing repositories must not be expected to allow for queries of labels other than those present in the ontology.

Annotation guidance

Note that this section is only a parking lot, the respective text will be moved into the AIRR Docs in the final version.

• Cells that come from Ficoll gradients should not be annotated as PBMCs as this is a sister node of lymphocyte. For the other sampling related fields, in nearly all cases venous blood (UBERON:0013756) will be the correct tissue and it should be used in the case of sample_type:peripheral venous puncture. However, if the mode of sampling is not specified, blood (UBERON:0000178) should be used instead. Also see https: //github.com/airr-community/airr-standards/issues/242

2.3.6 Schema Release Notes

Version 1.3.1: October 13, 2020

Version 1.3 documentation patch release.

Alignment Schema:

1. Added the deprecation tags for rearrangement_id, which were accidentally left out of the v1.3.0 release.

Version 1.3.0: May 28, 2020

Version 1.3 schema release.

New Schema:

- 1. Introduced the Repertoire Schema for describing study meta data.
- 2. Introduced the PCRTarget Schema for describing primer target locations.
- 3. Introduced the SampleProcessing Schema for describing experimental processing steps for a sample.
- 4. Replaced the SoftwareProcessing schema with the DataProcessing schema.
- 5. Introduced experimental schema for clonal clusters, lineage trees, tree nodes, and cells as Clone, Tree, Node, and Cell objects, respectively.

General Updates:

- 1. Added multiple additional attributes to a large number of schema propertes as AIRR extension attributes in the x-airr field. The new Attributes object contains definitions for these x-airr field attributes.
- 2. Added the top level required property to all relevant schema objects.
- 3. Added the title attribute containing the short, descriptive name to all relevant schema object fields.

4. Added an example attribute containing an example data value to multiple schema object fields.

AIRR Data Commons API:

1. Added OpenAPI V2 specification (specs/adc-api.yaml) for AIRR Data Commons API major version 1.

Ontology Support:

- 1. Added Ontology and CURIEResolution objects to support ontologies.
- 2. Added vocabularies/ontologies as JSON string for: Cell subset, Target substrate, Library generation method, Complete sequences, Physical linkage of different loci.

Rearrangement Schema:

- 1. Added the complete_vdj field to annotate whether a V(D)J alignment was full length.
- 2. Added the junction_length_aa field defining the length of the junction amino acid sequence.
- 3. Added the repertoire_id, sample_processing_id, and data_processing_id fields to serve as linkers to the appropriate metadata objects.
- 4. Added a controlled vocabulary to the locus field: IGH, IGI, IGK, IGL, TRA, TRB, TRD, TRG.
- 5. Deprecated the rearrangement_set_id and germline_database fields.
- 6. Deprecated rearrangement_id field and made the sequence_id field be the primary unique identifer for a rearrangement record, both in files and data repositories.
- 7. Added support secondary D gene rearrangement through the additional fields: d2_call, d2_score, d2_identity, d2_support, d2_cigar np3, np3_aa, np3_length, n3_length, p5d2_length, p3d2_length, d2_sequence_start, d2_sequence_end, d2_germline_start, d2_germline_start, d2_alignment_start, d2_alignment_end, d2_sequence_alignment, d2_sequence_alignment_aa, d2_germline_alignment_aa.
- 8. Updated field definitions with more concise V(D)J call descriptions.

Alignment Schema:

- 1. Deprecated the rearrangement_set_id and germline_database fields.
- 2. Added the data_processing_id field.

Study Schema:

1. Added the study_type field containing an ontology defined term for the study design.

Subject Schema:

- 1. Deprecated the organism field in favor of the new species field.
- 2. Deprecated the age field.
- 3. Introduced age ranges: age_min, age_max, and age_unit.

Diagnosis Schema:

1. Changed the type of the disease_diagnosis field from string to Ontology.

Sample Schema:

1. Changed the type of the tissue field from string to Ontology.

CellProcessing Schema:

- 1. Changed the type of the cell_subset field from string to Ontology.
- 2. Introduced the cell_species field which denotes the species from which the analyzed cells originate.

NucleicAcidProcessing Schema:

- 1. Defined the template_class field as type string.
- 2. Added a controlled vocabulary the library_generation_method field.
- 3. Changed the controlled vocabulary terms of complete_sequences. Replacing complete & untemplated with complete+untemplated and adding mixed.
- 4. Added the pcr_target field referencing the new PCRTarget schema object.

SequencingRun Schema:

- 1. Added the sequencing_run_id field which serves as the object identifer field.
- 2. Added the sequencing_files field which links to the RawSequenceData schema objects defining the raw read data.

RawSequenceData Schema:

- 1. Added the file_type field defining the sequence file type. This field is a controlled vocabulary restricted to: fasta, fastq.
- 2. Added the paired_read_length field defining mate-pair read lengths.
- 3. Defined the read_direction and paired_read_direction fields as type string.

DataProcessing Schema:

- 1. Replaces the SoftwareProcessing object.
- 2. Added data_processing_id, primary_annotation, data_processing_files, germline_database and analysis_provenance_id fields.

Version 1.2.1: Oct 5, 2018

Minor patch release.

- 1. Schema gene vs segment terminology corrections
- 2. Added Info object
- 3. Updated cell_subset URL in AIRR schema

Version 1.2.0: Aug 18, 2018

Peer reviewed released of the Rearrangement schema.

- 1. Definition change for the coordinate fields of the Rearrangement and Alignment schema. Coordinates are now defined as 1-based closed intervals, instead of 0-based half-open intervals (as previously defined in v1.1 of the schema).
- 2. Removed foreign study_id fields
- 3. Introduced keywords_study field

Version 1.1.0: May 3, 2018

Initial public released of the Rearrangement and Alignment schemas.

- 1. Added required and nullable constrains to AIRR schema.
- 2. Schema definitions for MiAIRR attributes and ontology.

- 3. Introduction of an x-airr object indicating if field is required by MiAIRR.
- 4. Rename rearrangement_set_id to data_processing_id.
- 5. Rename study_description to study_type.
- 6. Added physical_quantity format.
- 7. Raw sequencing files into separate schema object.
- 8. Rename Attributes object.
- 9. Added primary_annotation and repertoire_id.
- 10. Added diagnosis to repertoire object.
- 11. Added ontology for organism.
- 12. Added more detailed specification of sequencing_run, repertoire and rearrangement.
- 13. Added repertoire schema.
- 14. Rename definitions.yaml to airr-schema.yaml.
- 15. Removed c_call, c_score and c_cigar from required as this is not typical reference aligner output.
- 16. Renamed vdj_score, vdj_identity, vdj_evalue, and vdj_cigar to score, identity, evalue, and cigar.
- 17. Added missing c_identity and c_evalue fields to Rearrangement spec.
- 18. Swapped order of N and S operators in CIGAR string.
- 19. Some description clean up for consistency in Rearrangement spec.
- 20. Remove repeated objects in definitions.yaml.
- 21. Added Alignment object to definitions.yaml.
- 22. Updated MiARR format consistency check TSV with junction change.
- 23. Changed definition from functional to productive.

Version 1.0.1: Jan 9, 2018

MiAIRR v1 official release and initial draft of Rearrangement and Alignment schemas.

2.4 Data Submission and Query

2.4.1 Data Submission Guides for AIRR-seq studies

There are multiple data repositories that accept submission of AIRR-seq datasets. Each provides different capabilities but all comply with the MiAIRR standard.

CAIRR Pipeline

Introduction: The CAIRR pipeline for submitting standards-compliant B and T cell receptor repertoire sequencing studies to the NCBI

AIRR sequencing (AIRR-seq) has tremendous potential to understand the dynamics of the immune repertoire in vaccinology, infectious disease, autoimmunity, and cancer biology. The adaptation of high-throughput sequencing (HTS) for AIRR (Adaptive Immune Receptor Repertoire) studies has made possible to characterize the AIRR at unprecedented depth and the outcome of such sequencing produces big data. Effective sharing of AIRR-seq big data could potentially reveal amazing scientific insights. The AIRR Community has proposed MiAIRR (Minimum information about an Adaptive Immune Receptor Repertoire Sequencing Experiment), a standard for reporting AIRR-seq studies. The MiAIRR standard has been implemented using the National Center for Biotechnology Information (NCBI) repositories. Submissions of AIRR-seq data to the NCBI repositories typically use a combination of web-based and flat-file templates and include only a minimal amount of terminology validation. As a result, AIRR-seq studies at the NCBI are often described using inconsistent terminologies, limiting scientists' ability to access, find, interoperate, and reuse the data sets and to understand how the experiments were performed. CEDAR (Center for Expanded Data Annotation and Retrieval) develops technologies involving the use of data standards and ontologies to improve metadata quality. In order to improve metadata quality and ease AIRR-seq study submission process, we have developed an AIRR-seq data submission pipeline named CEDAR-AIRR (CAIRR). CAIRR leverages CEDAR's technologies to: i) create webbased templates whose entries are controlled by ontology terms, ii) generate and validate metadata and iii) submit the ontology-linked metadata and sequence files (FASTQ) to the NCBI BioProject, BioSample, and Sequence Read Archive (SRA) databases. Thus, CAIRR provides a web-based metadata submission interface that supports compliance with MiAIRR standards. The interface enables ontology-based validation for several data elements, including: organism, disease, cell type and subtype, and tissue. This pipeline will facilitate the NCBI submission process and improve the metadata quality of AIRR-seq studies.

Submission Steps

The submission steps are described in the MiAIRR-to-NCBI Submission Manual: *Option 1. Submission via the CEDAR system (CAIRR submission pipeline)*. You will need a CEDAR system account; you can self-register at https://cedar.metadatacenter.org. You will also need the identifier of a BioProject already entered in the NCBI BioProject database.

Citing the MiAIRR Pipeline

Bukhari, Syed Ahmad Chan, Martin J. O'Connor, Marcos Martínez-Romero, Attila L. Egyedi, Debra Debra Willrett, John Graybeal, Mark A. Musen, Florian Rubelt, Kei H. Cheung, and Steven H. Kleinstein. The CAIRR pipeline for submitting standards-compliant B and T cell receptor repertoire sequencing studies to the NCBI. Frontiers in Immunology 9 (2018): 1877. DOI: 10.3389/fimmu.2018.01877

Tell Us About It

Please let us know how it went! If you are willing, we would love to have your comments in a short survey, it should just take 5 minutes or so.

We also welcome entry of issues and requests in our github repository issues, and emails can be sent to cedarusers@lists.stanford.edu. Both of these resources are publicly visible.

Support or Contact

Having trouble with NCBI submission process through our pipeline? Please email to Syed Ahmad Chan Bukhari or to Marcos Martínez-Romero and we will help you sort it out.

Introduction to VDJServer

VDJServer is a cloud-based analysis portal for immune repertoire sequence data that provides access to a suite of tools for a complete analysis workflow, including modules for preprocessing and quality control of sequence reads, V(D)J gene assignment, repertoire characterization, and repertoire comparison. VDJServer also provides sophisticated visualizations for exploratory analysis. It is accessible through a standard web browser via a graphical user interface designed for use by immunologists, clinicians, and bioinformatics researchers. VDJServer provides a data commons for public sharing of repertoire sequencing data, as well as private sharing of data between users.

- VDJServer website
- VDJServer Community Data Portal
- Email VDJServer for information about submission of your study data.

References

Introduction to iReceptor

iReceptor is a platform for storing, sharing, and exploring AIRR-seq data according to the AIRR Community standards.

- iReceptor Website (General information)
- iReceptor Gateway (AIRR Data Commons data query and federation)
- iReceptor Repositories (AIRR Data Commons repositories)
- iReceptor Turnkey GitHUb (Software)
- Email iReceptor (Contact).

References

2.4.2 Data Submission for Inferred Genes and Alleles

In 2017, The AIRR Community established the Inferred Allele Review Committee (IARC) to evaluate inferred alleles for inclusion in relevant germline databases. IARC has worked, together with colleagues at IMGT and the US National Institutes of Health, to establish a systematic submission and review process. OGRDB was created and designed to support that process, and provide a real-time record of affirmed sequences.

OGRDB - reference database of inferred immune receptor genes

In recent years it has become possible to sequence immune receptor repertoires (immunoglobulins and T cell receptors) at great depth. The accurate analysis of these repertoires requires a comprehensive understanding of the germline genes that give rise to the repertoire through V(D)J gene recombination.

Even for well-studied species such as humans and mice, our knowledge of allelic variation is incomplete. Identifying new immunoglobulin and T cell receptor polymorphisms from the genome using traditional methods is technically challenging, because of the complex sequence architecture and repetitive nature of these loci. More recently, methods have been developed to infer novel sequences and alleles from sequenced repertoires.

The Adaptive Immune Receptor Repertoire (AIRR) Community was formed to promote and share good practice in adaptive immune repertoire sequencing. In 2017, it established the Inferred Allele Review Committee (IARC) to evaluate inferred alleles for inclusion in relevant germline databases. IARC's work is outlined in more detail in a poster, which was presented at a Systems Immunology Workshop at the University of Surrey, England, in March

2018, and in a recent paper. IARC has worked, together with colleagues at IMGT and the US National Institute of Health, to establish a systematic submission and review process. OGRDB was created and designed to support that process, and provide a real-time record of affirmed sequences. Affirmed sequences will be listed under the Sequences tab above, and the submissions that underpin them will be found under the Submissions tab. You can make your own submissions by following the steps below.

How to submit your sequences

As a first step, IARC is now ready to review submissions of inferred human IGHV genes and alleles. These sequences may be novel, or may extend incomplete sequences currently in the IMGT reference directories. Researchers interested in submitting sequences should:

Submission of IARC gene inference data to NCBI

General outline

IARC submission currently follows a "INSDC first" approach, means that all sequence data related to the reported inference is REQUIRED to be properly deposited in a general purpose sequence repository before it is reviewed by IARC. The submitter needs to complete the initial steps of submission to one of the INSDC repositories. Upon submission to IARC, some of this data will be pulled in from NCBI (TODO: What kind of data can we actually pull down from INSDC?)

The aim of this procedure is to ensure that inferences reviewed by IARC are public and will remain available in the long run. It is however explicitly *not* the aim to provide data that deterministically will yield the same inference results.

Deposition of inferred gene data at NCBI

At the end of the deposition process there should be three types of records present at NCBI:

- 1. A single record containing the final and full-length inferred sequence. The record is deposited in one of the following:
 - Genbank: All inferences that have been performed on the submitters own data CAN be submitted as [???] to Genbank. Note that Genbank typically only holds data that has a physical correlate which is not necessarily true for inferred sequences. Nevertheless NCBI currently accepts this as a kind of consensus building if it is performed on your own data. The Genbank record MUST link to the select set record (see 3.) via the DBLINK/DR field. Genbank records will be publicly available independent of other publications. Note that the for Genbank, the DBLINK field does not appear to be available through the BankIt submission interface. You can use Tbl2asn and Sequin, and edit the DBLINK field manually (as "Sequence Read Archive" is not one of the options on the template creation page. A sample Genbank deposit can be found under accession MK321694.
 - TPA (Third-party annotation): A segment of Genbank dedicated inferences. Also the TPA record MUST link to the select set record (see 3.) via the DBLINK/DR field. Note that in contrast to Genbank, TPA does REQUIRE a peer-reviewed publication describing the details of the inference process before the record will be made publicly available. A sample TPA deposit can be found under accession BK01573.

The format for both record types the Genbank format (link) with a standardized feature table (FT). Note that your initial submission MUST NOT contain any potential name for the gene as this will be assigned by IARC later on.

TODO: Is there any metadata that should be provided into the GB record?

- 2. One or multiple SRA records containing all raw reads of the the respective sequencing run. Note that if you are performing inference using third-party data, these records MUST be submitted by the original owner of the data. These record type will typically be present before the other. The metadata annotation of the records SHOULD be MiAIRR compliant [Rubelt et al.].
- 3. One or multiple SRA records containing the select set of reads from (2). The aim of these records is to document the number, quality, coverage and diversity of the reads in a dataset that _potentially_ support the inference. This means that the select set SHOULD be a superset of the reads that support the inference. It is NOT REQUIRED that inference tools deterministically return the inferred allele upon being fed with the select set. Generation of the select set from the complete set is described below. When submitting the select set to SRA the metadata context, i.e. the original links to project, sample and (if possible) experiment) SHOULD be maintained. Reads originating from multiple subjects or samples MUST NOT be pooled into a single new entry. The new record SHOULD be titled "Reads from <original_run_accession> supporting inference of Homo sapiens immunoglobulin heavy chain variable gene" and contain a design description, e.g., "Experimental workflow as described in original SRA/ENA record [<run_accession>]. Gene inference was performed using <software+version+parameters>. The reported reads were selected based on <selection_criteria>."

NOTE: It is reasonably likely, in the short term, that you will encounter questions from the SRA/ENA/Genbank staff about the nature of these deposits. If so, you can respond that they are made as part of a community effort to document novel alleles with an emphasis on transparency in data provenance. You can link to the IARC page and note that we worked together with IMGT and Genbank/TPA staff in designing this procedure.

Generating the select set

Below is the current procedure describing how to generate a select set using general purpose tools. This procedure was designed in a rather generic fashion so that it is easy to implement and does NOT REQUIRE inference tools to provide their own mechanisms. Note that it is currently assumed that the procedure is not fully deterministic, i.e. the select set cannot simply be generated using the complete read data and the inferred sequence, as there are additional filter criteria that apply. In addition the select set SHOULD not be subject to any modifications that are not listed below. This includes UMI-based consensus building or other aggregation steps that are not fully transparent to a third-party.

- 1. Assemble paired-end reads. The two reads MUST overlap. Recommended tool: PandaSeq
- 2. Perform PHRED filtering that is equivalent to the one performed by inference pipeline. Recommended tool: Immcantation suite
- 3. Perform a *blastn* search using the data from (2.) as query and bp 1-312 of the inferred gene as reference library. Require matches to be full-length and >99.6% ID. Record all matching read ID. Recommended tool: NCBI BLAST
- 4. Select the reads with the read ID found in (3.) from the original unmerged FASTQs. Note that each select set MUST be derived from a single donor and sample. Recommended tool: Christian's cryptic extractor script
- 5. Submit the select set to SRA

Submit the inferred sequences to IARC via OGRDB, following the

• OGRDB Submission Guide

Additional information is available at the

• OGRDB Website

References

2.4.3 Data Query and Download from the AIRR Data Commons

Submission of AIRR-seq datasets to public data repositories means that other researchers can query, download and reuse that data for novel analyses.

AIRR Data Commons

The AIRR Data Commons is a network of distributed repositories that store AIRR-seq data and adhere to the AIRR Community standards. We define the AIRR Data Commons as consisting of the set of repositories that both:

- Adhere to the AIRR Common Repositories Working Group recommendations for promoting, sharing, and use of AIRR-seq data.
- Implement the ADC API as a programmatic mechanism to access that data.

More information on repositories in the AIRR Data Commons and how to query these repositories can be found on the AIRR Data Commons page:

AIRR Data Commons

The use of high-throughput sequencing for profiling B-cell and T-cell receptors has resulted in a rapid increase in data generation. It is timely, therefore, for the Adaptive Immune Receptor Repertoire (AIRR) community to establish a clear set of community-accepted data and metadata standards; analytical tools; and policies and practices for infrastructure to support data deposit, curation, storage, and use. Such actions are in accordance with international funder and journal policies that promote data deposition and data sharing – at a minimum, data on which scientific publications are based should be made available immediately on publication. Data deposit in publicly accessible databases ensures that published results may be validated. Such deposition also facilitates reuse of data for the generation of new hypotheses and new knowledge.

The AIRR Common Repository Working Group (CRWG) has developed a set of recommendations that promote the deposit, sharing, and use of AIRR sequence data. These recommendations were refined following community discussions at the AIRR 2016 and 2017 Community Meetings and were approved through a vote by the AIRR Community at the AIRR Community Meeting in December 2017. Updates to these recommendations have continued, with the latest set of Recommendations ratified at the AIRR Community meeting in May 2019.

In May 2020, the AIRR Community released the first verion of the AIRR Data Commons Application Programming Interface (ADC API), a specification for programmatic access to query and download AIRR-seq data from repositories that adhere to the AIRR Standards. We define the AIRR Data Commons as consisting of the set of repositories that:

- · adhere to the CRWG recommendations for promoting, sharing, and use of AIRR-seq data, and
- that implement the ADC API as a programmatic mechanism to access that data.

This page provides a central location for the community to discover resources that belong to the AIRR Data Commons.

AIRR Data Commons Repositories

These data repositories all implement the AIRR Data Commons (ADC) API programmatic access to query and down-load AIRR-seq data.

- *iReceptor Public Archive*
- VDJServer Community Data Portal

Querying the AIRR Data Commons

Each of the repositories above can be queried directly using the *ADC API*. In addition, the following tools and platforms implement web based user interfaces that use the ADC API to query repositories in the AIRR Data Commons:

• *iReceptor Gateway*

There are *query and analysis use cases* and *a set of example queries* available for the AIRR Data Commons and the ADC API.

Other Public AIRR-Seq Repositories

There are additional data repositories that provide access to AIRR-seq data but which did not implement the ADC API for programmatic access. Information about some of these repositories are provided in a B-T.CR forum post.

Germline Gene Inference and Usage

- *OGRDB* provides a list of alleles affirmed by the AIRR Community's Inferred Allele Review Committee, together with supporting information.
- VDJbase provides gene usage information derived from a growing base of AIRR-seq repertoires, including inferred genotypes and haplotypes.

2.5 Software

2.5.1 AIRR Standards Reference Implementations

AIRR Python Reference Library

The airr reference library provides basic functions and classes for interacting with AIRR Community Data Representation Standards, including tools for read, write and validation.

API Reference

Rearrangement Interface

airr.**read_rearrangement** (*filename*, *validate=False*, *debug=False*) Open an iterator to read an AIRR rearrangements file

Parameters

- **file** (*str*) path to the input file.
- **validate** (*bool*) whether to validate data as it is read, raising a ValidationError exception in the event of an error.
- debug (bool) debug flag. If True print debugging information to standard error.

Returns iterable reader class.

Return type *airr.io.RearrangementReader*

airr.create_rearrangement (filename, fields=None, debug=False)

Parameters

- **filename** (*str*) output file path.
- **fields** (*list*) additional non-required fields to add to the output.
- **debug** (bool) debug flag. If True print debugging information to standard error.

Returns open writer class.

Return type *airr.io.RearrangementWriter*

airr.derive_rearrangement (*out_filename*, *in_filename*, *fields=None*, *debug=False*) Create an empty AIRR rearrangements file with fields derived from an existing file

Parameters

- **out_filename** (*str*) output file path.
- **in_filename** (*str*) existing file to derive fields from.
- **fields** (*list*) additional non-required fields to add to the output.
- **debug** (bool) debug flag. If True print debugging information to standard error.

Returns open writer class.

Return type *airr.io.RearrangementWriter*

airr.load_rearrangement (filename, validate=False, debug=False)

Load the contents of an AIRR rearrangements file into a data frame

Parameters

- **filename** (*str*) input file path.
- **validate** (*bool*) whether to validate data as it is read, raising a ValidationError exception in the event of an error.
- **debug** (bool) debug flag. If True print debugging information to standard error.

Returns Rearrangement records as rows of a data frame.

Return type pandas.DataFrame

airr.dump_rearrangement (dataframe, filename, debug=False)

Write the contents of a data frame to an AIRR rearrangements file

Parameters

- **dataframe** (*pandas*. *DataFrame*) **data frame of rearrangement data**.
- **filename** (*str*) output file path.
- **debug** (bool) debug flag. If True print debugging information to standard error.

Returns True if the file is written without error.

Return type bool

airr.merge_rearrangement (*out_filename*, *in_filenames*, *drop=False*, *debug=False*) Merge one or more AIRR rearrangements files

Parameters

- **out_filename** (*str*) output file path.
- **in_filenames** (*list*) list of input files to merge.

- **drop** (bool) drop flag. If True then drop fields that do not exist in all input files, otherwise combine fields from all input files.
- **debug** (bool) debug flag. If True print debugging information to standard error.

Returns True if files were successfully merged, otherwise False.

Return type bool

airr.validate_rearrangement (filename, debug=False) Validates an AIRR rearrangements file

Parameters

- **filename** (*str*) path of the file to validate.
- debug (bool) debug flag. If True print debugging information to standard error.

Returns True if files passed validation, otherwise False.

Return type bool

Repertoire Interface

```
airr.load_repertoire (filename, validate=False, debug=False)
```

Load an AIRR repertoire metadata file

Parameters

- **filename** (*str*) path to the input file.
- **validate** (bool) whether to validate data as it is read, raising a ValidationError exception in the event of an error.
- **debug** (bool) debug flag. If True print debugging information to standard error.

Returns list of Repertoire dictionaries.

Return type list

```
airr.write_repertoire (filename, repertoires, info=None, debug=False)
Write an AIRR repertoire metadata file
```

Parameters

- file (str) path to the output file.
- **repertoires** (*list*) array of repertoire objects.
- **info** (*object*) info object to write. Will write current AIRR Schema info if not specified.
- **debug** (bool) debug flag. If True print debugging information to standard error.

Returns True if the file is written without error.

Return type bool

airr.validate_repertoire (filename, debug=False)

Validates an AIRR repertoire metadata file

Parameters

- **filename** (*str*) path of the file to validate.
- **debug** (bool) debug flag. If True print debugging information to standard error.

Returns True if files passed validation, otherwise False.

Return type bool

airr.repertoire_template()

Return a blank repertoire object from the template. This object has the complete structure with all of the fields and all values set to None or empty string.

Returns empty repertoire object.

Return type object

Classes

class airr.io.**RearrangementReader** (*handle*, *base=1*, *validate=False*, *debug=False*) Iterator for reading Rearrangement objects in TSV format

fields

field names in the input Rearrangement file.

Type list

external_fields

list of fields in the input file that are not part of the Rearrangement definition.

Type list

____init___ (handle, base=1, validate=False, debug=False) Initialization

Parameters

- handle (file) file handle of the open Rearrangement file.
- **base** (*int*) one of 0 or 1 specifying the coordinate schema in the input file. If 1, then the file is assumed to contain 1-based closed intervals that will be converted to python style 0-based half-open intervals for known fields. If 0, then values will be unchanged.
- **validate** (bool) perform validation. If True then basic validation will be performed will reading the data. A ValidationError exception will be raised if an error is found.
- **debug** (bool) debug state. If True prints debug information.

Returns reader object.

Return type airr.io.RearrangementReader

___iter__()

Iterator initializer

Returns airr.io.RearrangementReader

__next__()

Next method

Returns parsed Rearrangement data.

Return type dict

close()

Closes the Rearrangement file

next()

Next method

```
class airr.io.RearrangementWriter (handle, fields=None, base=1, debug=False)
Writer class for Rearrangement objects in TSV format
```

fields

field names in the output Rearrangement file.

Type list

external_fields

list of fields in the output file that are not part of the Rearrangement definition.

Type list

___init___(handle, fields=None, base=1, debug=False)

Initialization

Parameters

- handle (file) file handle of the open Rearrangements file.
- **fields** (*list*) list of non-required fields to add. May include fields undefined by the schema.
- **base** (*int*) one of 0 or 1 specifying the coordinate schema in the output file. Data provided to the write is assumed to be in python style 0-based half-open intervals. If 1, then data will be converted to 1-based closed intervals for known fields before writing. If 0, then values will be unchanged.
- **debug** (*bool*) debug state. If True prints debug information.

Returns writer object.

Return type *airr.io.RearrangementWriter*

close()

Closes the Rearrangement file

write(row)

Write a row to the Rearrangement file

Parameters row (*dict*) - row to write.

class airr.schema.Schema (*definition*) AIRR schema definitions

properties

field definitions.

Type collections.OrderedDict

info

schema info.

Type collections.OrderedDict

required

list of mandatory fields.

Type list

optional

list of non-required fields.

Type list

false_values

accepted string values for False.

Type list

true_values

accepted values for True.

Type list

from_bool (*value*, *validate=False*) Converts a boolean to a string

Parameters

- **value** (*bool*) logical value.
- **validate** (*bool*) when True raise a ValidationError for an invalid value. Otherwise, set invalid values to None.

Returns conversion of True or False or 'T' or 'F'.

Return type str

Raises airr.ValidationError - raised if value is invalid when validate is set True.

spec (field)

Get the properties for a field

Parameters name (*str*) – field name.

Returns definition for the field.

Return type collections.OrderedDict

to_bool (value, validate=False)

Convert a string to a boolean

Parameters

- **value** (*str*) logical value as a string.
- **validate** (*bool*) when True raise a ValidationError for an invalid value. Otherwise, set invalid values to None.

Returns conversion of the string to True or False.

Return type bool

Raises airr.ValidationError - raised if value is invalid when validate is set True.

to_float (value, validate=False)

Converts a string to a float

Parameters

- **value** (*str*) float value as a string.
- **validate** (*bool*) when True raise a ValidationError for an invalid value. Otherwise, set invalid values to None.

Returns conversion of the string to a float.

Return type float

Raises airr.ValidationError - raised if value is invalid when validate is set True.

to_int (value, validate=False)

Converts a string to an integer

Parameters

- **value** (*str*) integer value as a string.
- **validate** (*bool*) when True raise a ValidationError for an invalid value. Otherwise, set invalid values to None.

Returns conversion of the string to an integer.

Return type int

Raises airr.ValidationError - raised if value is invalid when validate is set True.

type (field)

Get the type for a field

Parameters name (*str*) – field name.

Returns the type definition for the field

Return type str

validate_header(header)

Validate header against the schema

Parameters header (*list*) – list of header fields.

Returns True if a ValidationError exception is not raised.

Return type bool

Raises airr.ValidationError - raised if header fails validation.

validate_object (obj, missing=True, nonairr=True, context=None)
Validate Repertoire object data against schema

Parameters

- **obj** (*dict*) dictionary containing a single repertoire object.
- **missing** (*bool*) provides warnings for missing optional fields.
- (bool (nonairr) provides warning for non-AIRR fields that cannot be validated.
- **context** (*string*) used by recursion to indicate place in object hierarchy

Returns True if a ValidationError exception is not raised.

Return type bool

Raises airr.ValidationError – raised if object fails validation.

validate_row(row)

Validate Rearrangements row data against schema

Parameters row (*dict*) – dictionary containing a single record.

Returns True if a ValidationError exception is not raised.

Return type bool

Raises airr.ValidationError - raised if row fails validation.

Schema

airr.schema.AlignmentSchema Schema object for the Alignment definition AIRR schema definitions

airr.schema.**properties** field definitions.

Type collections.OrderedDict

airr.schema.info schema info.

Type collections.OrderedDict

airr.schema.**required** list of mandatory fields.

Type list

airr.schema.**optional** list of non-required fields.

Type list

airr.schema.false_values accepted string values for False.

Type list

airr.schema.true_values accepted values for True.

Type list

airr.schema.RearrangementSchema Schema object for the Rearrangement definition AIRR schema definitions

airr.schema.properties field definitions.

Type collections.OrderedDict

airr.schema.**info** schema info.

Type collections.OrderedDict

airr.schema.**required** list of mandatory fields.

Type list

airr.schema.**optional** list of non-required fields.

Type list

airr.schema.false_values accepted string values for False.

Type list

airr.schema.true_values accepted values for True.

Type list

airr.schema.RepertoireSchema Schema object for the Repertoire definition AIRR schema definitions

airr.schema.properties field definitions.

Type collections.OrderedDict

airr.schema.**info** schema info.

Type collections.OrderedDict

airr.schema.**required** list of mandatory fields.

Type list

airr.schema.**optional** list of non-required fields.

Type list

airr.schema.**false_values** accepted string values for False.

Type list

airr.schema.true_values accepted values for True.

Type list

Commandline Tools

airr-tools

AIRR Community Standards utility commands.

usage: airr-tools [-h] [--version] ...

-h, --help

show this help message and exit

--version

show program's version number and exit

airr-tools merge

Merge AIRR rearrangement files.

```
usage: airr-tools merge [--version] [-h] -o OUT_FILE [--drop] -a AIRR_FILES [AIRR_FILES ...]
```

--version

show program's version number and exit

-h, --help

show this help message and exit

-o <out_file>
Output file name.

--drop

If specified, drop fields that do not exist in all input files. Otherwise, include all columns in all files and fill missing data with empty strings.

-a <airr_files> A list of AIRR rearrangement files.

A list of AIKK realitaligement life

airr-tools validate

```
Validate AIRR files.
```

usage: airr-tools validate [--version] [-h] ...

--version

show program's version number and exit

-h, --help

show this help message and exit

airr-tools validate rearrangement

Validate AIRR rearrangement files.

```
usage: airr-tools validate rearrangement [--version] [-h] -a AIRR_FILES [AIRR_FILES ...]
```

--version

show program's version number and exit

- -h, --help show this help message and exit
- **-a** <airr_files>

A list of AIRR rearrangement files.

airr-tools validate repertoire

Validate AIRR repertoire metadata files.

```
usage: airr-tools validate repertoire [--version] [-h] -a AIRR_FILES
[AIRR_FILES ...]
```

--version

show program's version number and exit

-h, --help

show this help message and exit

```
-a <airr_files>
```

A list of AIRR repertoire metadata files.

Python Library Release Notes

Version 1.3.1: October 13, 2020

- 1. Refactored merge_rearrangement to allow for larger number of files.
- 2. Improved error handling in format validation operations.

Version 1.3.0: May 30, 2020

- 1. Updated schema set to v1.3.
- 2. Added load_repertoire, write_repertoire, and validate_repertoire to airr. interface to read, write and validate Repertoire metadata, respectively.
- 3. Added repertoire_template to airr.interface which will return a complete repertoire object where all fields have null values.
- 4. Added validate_object to airr.schema that will validate a single repertoire object against the schema.
- 5. Extended the airr-tools commandline program to validate both rearrangement and repertoire files.

Version 1.2.1: October 5, 2018

1. Fixed a bug in the python reference library causing start coordinate values to be empty in some cases when writing data.

Version 1.2.0: August 17, 2018

- 1. Updated schema set to v1.2.
- 2. Several improvements to the validate_rearrangement function.
- 3. Changed behavior of all *airr.interface* functions to accept a file path (string) to a single Rearrangement TSV, instead of requiring a file handle as input.
- 4. Added base argument to RearrangementReader and RearrangementWriter to support optional conversion of 1-based closed intervals in the TSV to python-style 0-based half-open intervals. Defaults to conversion.
- 5. Added the custom exception ValidationError for handling validation checks.
- 6. Added the validate argument to RearrangementReader which will raise a ValidationError exception when reading files with missing required fields or invalid values for known field types.
- 7. Added validate argument to all type conversion methods in Schema, which will now raise a ValidationError exception for value that cannot be converted when set to True. When set False (default), the previous behavior of assigning None as the converted value is retained.
- Added validate_header and validate_row methods to Schema and removed validations methods from RearrangementReader.
- 9. Removed automatic closure of file handle upon reaching the iterator end in RearrangementReader.

Version 1.1.0: May 1, 2018

Initial release.

Installation

Install in the usual manner from PyPI:

```
> pip3 install airr --user
```

Or from the downloaded source code directory:

```
> python3 setup.py install --user
```

Quick Start

Reading AIRR Repertoire metadata files

The airr package contains functions to read and write AIRR repertoire metadata files. The file format is either YAML or JSON, and the package provides a light wrapper over the standard parsers. The file needs a json, yaml, or yml file extension so that the proper parser is utilized. All of the repertoires are loaded into memory at once and no streaming interface is provided:

```
import airr
# Load the repertoires
data = airr.load_repertoire('input.airr.json')
for rep in data['Repertoire']:
    print(rep)
```

Why are the repertoires in a list versus in a dictionary keyed by the repertoire_id? There are two primary reasons for this. First, the repertoire_id might not have been assigned yet. Some systems might allow MiAIRR metadata to be entered but the repertoire_id is assigned to that data later by another process. Without the repertoire_id, the data could not be stored in a dictionary. Secondly, the list allows the repertoire data to have a default ordering. If you know that the repertoires all have a unique repertoire_id then you can quickly create a dictionary object using a comprehension:

```
rep_dict = { obj['repertoire_id'] : obj for obj in data['Repertoire'] }
```

Writing AIRR Repertoire metadata files

Writing AIRR repertoire metadata is also a light wrapper over standard YAML or JSON parsers. The airr library provides a function to create a blank repertoire object in the appropriate format with all of the required fields. As with the load function, the complete list of repertoires are written at once, there is no streaming interface:

```
import airr
# Create some blank repertoire objects in a list
reps = []
for i in range(5):
    reps.append(airr.repertoire_template())
```

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```
# Write the repertoires
airr.write_repertoire('output.airr.json', reps)
```

Reading AIRR Rearrangement TSV files

The airr package contains functions to read and write AIRR rearrangement files as either iterables or pandas data frames. The usage is straightforward, as the file format is a typical tab delimited file, but the package performs some additional validation and type conversion beyond using a standard CSV reader:

```
import airr
# Create an iteratable that returns a dictionary for each row
reader = airr.read_rearrangement('input.tsv')
for row in reader: print(row)
# Load the entire file into a pandas data frame
df = airr.load_rearrangement('input.tsv')
```

Writing AIRR formatted files

Similar to the read operations, write functions are provided for either creating a writer class to perform row-wise output or writing the entire contents of a pandas data frame to a file. Again, usage is straightforward with the airr output functions simply performing some type conversion and field ordering operations:

```
import airr
# Create a writer class for iterative row output
writer = airr.create_rearrangement('output.tsv')
for row in reader: writer.write(row)
# Write an entire pandas data frame to a file
airr.dump_rearrangement(df, 'file.tsv')
```

By default, create_rearrangement will only write the required fields in the output file. Additional fields can be included in the output file by providing the fields parameter with an array of additional field names:

```
# Specify additional fields in the output
fields = ['new_calc', 'another_field']
writer = airr.create_rearrangement('output.tsv', fields=fields)
```

A common operation is to read an AIRR rearrangement file, and then write an AIRR rearrangement file with additional fields in it while keeping all of the existing fields from the original file. The derive_rearrangement function provides this capability:

```
import airr
# Read rearrangement data and write new file with additional fields
reader = airr.read_rearrangement('input.tsv')
fields = ['new_calc']
writer = airr.derive_rearrangement('output.tsv', 'input.tsv', fields=fields)
for row in reader:
```

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```
row['new_calc'] = 'a value'
writer.write(row)
```

Validating AIRR data files

The airr package can validate repertoire and rearrangement data files to insure that they contain all required fields and that the fields types match the AIRR Schema. This can be done using the airr-tools command line program or the validate functions in the library can be called:

```
# Validate a rearrangement file
airr-tools validate rearrangement -a input.tsv
# Validate a repertoire metadata file
airr-tools validate repertoire -a input.airr.json
```

Combining Repertoire metadata and Rearrangement files

The airr package does not keep track of which repertoire metadata files are associated with rearrangement files, so users will need to handle those associations themselves. However, in the data, the repertoire_id field forms the link. The typical usage is that a program is going to perform some computation on the rearrangements, and it needs access to the repertoire metadata as part of the computation logic. This example code shows the basic framework for doing that, in this case doing gender specific computation:

```
import airr
# Load the repertoires
data = airr.load_repertoire('input.airr.json')
# Put repertoires in dictionary keyed by repertoire_id
rep_dict = { obj['repertoire_id'] : obj for obj in data['Repertoire'] }
# Create an iteratable for rearrangement data
reader = airr.read_rearrangement('input.tsv')
for row in reader:
    # get repertoire metadata with this rearrangement
   rep = rep_dict[row['repertoire_id']]
    # check the gender
   if rep['subject']['sex'] == 'male':
        # do male specific computation
   elif rep['subject']['sex'] == 'female':
       # do female specific computation
    else:
        # do other specific computation
```

AIRR Data Representation Reference Library

airr is an R package for working with data formatted according to the AIRR Data Representation schemas. It includes the full set of schema definitions along with simple functions for read, write and validation.

Usage Vignette

Introduction

Since the use of High-throughput sequencing (HTS) was first introduced to analyze immunoglobulin (B-cell receptor, antibody) and T-cell receptor repertoires (Freeman et al, 2009; Robins et al, 2009; Weinstein et al, 2009), the increasing number of studies making use of this technique has produced enormous amounts of data and there exists a pressing need to develop and adopt common standards, protocols, and policies for generating and sharing data sets. The Adaptive Immune Receptor Repertoire (AIRR) Community formed in 2015 to address this challenge (Breden et al, 2017) and has stablished the set of minimal metadata elements (MiAIRR) required for describing published AIRR datasets (Rubelt et al, 2017) as well as file formats to represent this data in a machine-readable form. The airr R package provide read, write and validation of data following the AIRR Data Representation schemas. This vignette provides a set of simple use examples.

AIRR Data Representation Standards

The AIRR Community's recommendations for a minimal set of metadata that should be used to describe an AIRR-seq data set when published or deposited in a AIRR-compliant public repository are described in Rubelt et al, 2017. The primary aim of this effort is to make published AIRR datasets FAIR (findable, accessible, interoperable, reusable); with sufficient detail such that a person skilled in the art of AIRR sequencing and data analysis will be able to reproduce the experiment and data analyses that were performed.

Following this principles, V(D)J reference alignment annotations are saved in standard tab-delimited files (TSV) with associated metadata provided in accompanying YAML formatted files. The column names and field names in these files have been defined by the AIRR Data Representation Working Group using a controlled vocabulary of standardized terms and types to refer to each piece of information.

Reading AIRR formatted files

The airr package contains the function read_rearrangement to read and validate files containing AIRR Rearrangement records, where a Rearrangement record describes the collection of optimal annotations on a single sequence that has undergone V(D)J reference alignment. The usage is straightforward, as the file format is a typical tabulated file. The argument that needs attention is base, with possible values "0" and "1". base denotes the starting index for positional fields in the input file. Positional fields are those that contain alignment coordinates and names ending in "_start" and "_end". If the input file is using 1-based closed intervals (R style), as defined by the standard, then positional fields will not be modified under the default setting of base="1". If the input file is using 0-based coordinates with half-open intervals (python style), then positional fields may be converted to 1-based closed intervals using the argument base="0"."

```
library(airr)
```

```
example_data <- system.file("extdata", "rearrangement-example.tsv.gz", package="airr")
basename(example_data)</pre>
```

```
## [1] "rearrangement-example.tsv.gz"
```

```
airr_rearrangement <- read_rearrangement(example_data)
class(airr_rearrangement)</pre>
```

```
## [1] "spec_tbl_df" "tbl_df" "tbl" "data.frame"
```

head(airr_rearrangement)

```
## # A tibble: 6 x 33
## sequence_id sequence rev_comp productive vj_in_frame stop_codon v_call d_call j_
⇔cigar

      ##
      <chr>
      <lgl><lgl><lgl></lgl>

      $\mathcal{s}$<chr>
      <chr>
      <chr</td>
      <chr>
      <chr</td>
      <chr</td>
      <chr>
      <chr</td>
      <chr<</td>
      <chr</td>
      <chr<<t
                                                                                                                 <chr> <chr>
                                                                                               FALSE
                                                                                                                     IGHV2...
→IGHD5... IGHJ4... IGHG ..... CAGATCACCTTGAAG... TGTGCAC...
→CAHSAGWLPD... 20S56N... 274S5N...
## 2 SRR765688.... NNNNNNN... FALSE
                                                           TRUE
                                                                                 TRUE
                                                                                                     FALSE
                                                                                                                       IGHV5...
→IGHD3... IGHJ6... IGHG .....GAGGTGCAGCTGGTG... TGTGCGA....
→CARHGLYGCD... 20S40N... 305S29...
## 3 SRR765688.... NNNNNNN... FALSE
                                                               TRUE
                                                                                  TRUE
                                                                                                     FALSE
                                                                                                                       IGHV7...
→IGHD3... IGHJ4... IGHG
                                          ..... TGTGCGA....
→CAREERRSSG... 20S33N... 293S13...
## 4 SRR765688.... NNNNNNN... FALSE
                                                               TRUE
                                                                                                                       IGHV7....
                                                                                 TRUE
                                                                                                     FALSE
→IGHD3... IGHJ6... IGHG ..... CAGGTGCAGCTGGTG... TGTGCGA....
→CAREGYYFDT... 20S33N... 290S9N...
## 5 SRR765688.... NNNNNNN... FALSE
                                                              TRUE
                                                                                 TRUE
                                                                                                     FALSE
                                                                                                                       IGHV7...
→IGHD1... IGHJ6... IGHG
                                         ..... CAGGTGCAGCTGGTG... TGTGCGA...
→CARDSGGMDVW 20S33N... 283S4N...
## 6 SRR765688.... NNNNNNN... FALSE FALSE
                                                                                  TRUE
                                                                                                     TRUE
                                                                                                                        IGHV2...
→IGHD2... IGHJ4... IGHA
                                          ..... TGTGTCC....
→CVLSRRLGDS... 20S56N... 273S12...
## # ... with 17 more variables: j_cigar <chr>, v_sequence_start <int>, v_sequence_
## # d_sequence_end <int>, d_germline_start <int>, d_germline_end <int>, j_sequence_
→start <int>, j_sequence_end <int>, j_germline_start <int>, j_germline_end <int>,
## #
         junction_length <int>, np1_length <int>, np2_length <int>, duplicate_count
→<int>
```

Writing AIRR formatted files

The airr package contains the function write_rearrangement to write Rearrangement records to the AIRR TSV format.

out_file <- file.path(tempdir(), "airr_out.tsv")
write_rearrangement(airr_rearrangement, out_file)</pre>

References

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sequencing data. Nat Immunol 18 (12): 1274-8.

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Reference Topics

read_airr

Read an AIRR TSV

Description

read_airr reads a TSV containing AIRR records.

Usage

read_airr(file, base = c("1", "0"), schema = RearrangementSchema, ...)

```
read_rearrangement(file, base = c("1", "0"), ...)
```

```
read_alignment(file, base = c("1", "0"), ...)
```

Arguments

file input file path.

base starting index for positional fields in the input file. If "1", then these fields will not be modified. If "0", then
fields ending in "_start" and "_end" are 0-based half-open intervals (python style) in the input file and will
be converted to 1-based closed-intervals (R style).

schema Schema object defining the output format.

... additional arguments to pass to read_delim.

Value

A data.frame of the TSV file with appropriate type and position conversion for fields defined in the specification.

Details

read_rearrangement reads an AIRR TSV containing Rearrangement data.

read_alignment reads an AIRR TSV containing Alignment data.

Examples

```
# Get path to the rearrangement-example file
file <- system.file("extdata", "rearrangement-example.tsv.gz", package="airr")
# Load data file
df <- read_rearrangement(file)</pre>
```

See also

See Schema for the AIRR schema object definition. See write_airr for writing AIRR data.

write_airr

Write an AIRR TSV

Description

write_airr writes a TSV containing AIRR formatted records.

Usage

```
write_airr(data, file, base = c("1", "0"), schema = RearrangementSchema, ...)
```

write_rearrangement(data, file, base = c("1", "0"), ...)

```
write_alignment(data, file, base = c("1", "0"), ...)
```

Arguments

data data.frame of Rearrangement data.

file output file name.

base starting index for positional fields in the output file. Fields in the input data are assumed to be 1-based closedintervals (R style). If "1", then these fields will not be modified. If "0", then fields ending in _start and _end will be converted to 0-based half-open intervals (python style) in the output file.

schema Schema object defining the output format.

... additional arguments to pass to write_delim.

Details

write_rearrangement writes a data.frame containing AIRR Rearrangement data to TSV.

write_alignment writes a data.frame containing AIRR Alignment data to TSV.

Examples

```
# Get path to the rearrangement-example file
file <- system.file("extdata", "rearrangement-example.tsv.gz", package="airr")
# Load data file
df <- read_rearrangement(file)
# Write a Rearrangement data file
outfile <- file.path(tempdir(), "output.tsv")
write_rearrangement(df, outfile)
```

See also

See Schema for the AIRR schema object definition. See read_airr for reading to AIRR files.

validate_airr

Validate AIRR data

Description

validate_airr validates compliance of the contents of a data.frame to the AIRR data standards.

Usage

validate_airr(data, schema = RearrangementSchema)

Arguments

data data.frame to validate.

schema Schema object defining the data standard.

Value

Returns TRUE if the input data is compliant and FALSE if not.

Examples

```
# Get path to the rearrangement-example file
file <- system.file("extdata", "rearrangement-example.tsv.gz", package="airr")
# Load data file
df <- read_rearrangement(file)</pre>
```

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```
# Validate a data.frame against the Rearrangement schema
validate_airr(df, schema=RearrangementSchema)
```

[1] TRUE

load_schema

Load a schema definition

Description

load_schema loads an AIRR object definition from the internal definition set.

Usage

load_schema(definition)

Arguments

definition name of the schema definition.

Value

A Schema object for the definition.

Details

Valid definitions include:

- "Rearrangement"
- "Alignment"
- "Study"
- "Subject"
- "Diagnosis"
- "Sample"
- "CellProcessing"
- "NucleicAcidProcessing"
- "RawSequenceData"
- "SoftwareProcessing"

Examples

```
# Load the Rearrangement definition
schema <- load_schema("Rearrangement")
# Load the Alignment definition
schema <- load_schema("Alignment")</pre>
```

See also

See Schema for the return object.

Schema-class

S4 class defining an AIRR standard schema

Description

Schema defines a common data structure for AIRR Data Representation standards.

Usage

"names"(x)			
------------	--	--	--

"["(x, i)

"\$"(x, name)

AlignmentSchema

RearrangementSchema

Arguments

x Schema object.

i field name.

name field name.

Format

A Schema object.

An object of class Schema of length 1.

An object of class Schema of length 1.

Details

The following predefined Schema objects are defined: AlignmentSchema: AIRR Alignment Schema. RearrangementSchema: AIRR Rearrangement Schema.

Slots

required character vector of required fields.
optional character vector of non-required fields.
properties list of field definitions.

info list schema information.

See also

See load_schema for loading a Schema from the definition set. See read_airr, write_airr and validate_airr schema operators.

ExampleData

Example AIRR data

Description

Example data files compliant with the the AIRR Data Representation standards.

Format

extdata/rearrangement-example.tsv.gz: Rearrangement TSV file.

Examples

```
# Get path to the rearrangement-example file
file <- system.file("extdata", "rearrangement-example.tsv.gz", package="airr")
# Load data file
df <- read_rearrangement(file)</pre>
```

R Library Release Notes

Version 1.3.0: May 26, 2020

1. Updated schema set to v1.3.

2. Added info slot to Schema object containing general schema information.

Version 1.2.0: August 17, 2018

- 1. Updated schema set to v1.2.
- 2. Changed defaults to base="1" for read and write functions.
- 3. Updated example TSV file with coordinate changes, addition of germline_alignment data and simplification of sequence_id values.

Version 1.1.0: May 1, 2018

Initial release.

Download & Installation

To install the latest release from CRAN:

install.packages("airr")

To build from the source code, first install the build dependencies:

```
install.packages(c("devtools", "roxygen2"))
```

To install the latest development code via devtools:

```
library(devtools)
install_github("airr-community/airr-standards/lang/R@master")
```

Note, using install_github will not build the documentation. To generate the documentation, clone the repository, and then build as normal using the following R commands from the package root lang/R:

```
library(devtools)
install_deps(dependencies=T)
document()
install()
```

Dependencies

Imports: methods, readr, stats, stringi, yaml **Suggests:** knitr, rmarkdown, testthat

Authors

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ADC API Reference Implementation

The AIRR Community provides a reference implementation for an ADC API service. The reference implementation can be utilized for any number of tasks. For example, a data repository might use the source code as a starting point for their own implementation and can compare the behaviour of their service against the reference. Another example is a tool developer, who wishes to use the API, can setup a local data repository so they can develop and test their tool before sending API requests across the internet to remote data repositories. While the reference implementation is functionally complete, it has minimal security and no optimizations for large data so it should not be used directly for production systems.

The reference implementation consists of three GitHub repositories: adc-api, adc-api-js-mongodb, and adc-apimongodb-repository. The three repositories correspond to the top-level service composition (adc-api), a JavaScript web service that responds to API requests and queries a MongoDB database (adc-api-js-mongodb), and a MongoDB database for holding AIRR-seq data (adc-api-mongodb-repository). Docker and docker-compose are used to provide a consistent deployment environment and compose the multiple components together into a single service. Complete documentation for configuring and deploying the reference implementation is available in the adc-api repository.

2.5.2 Resources and Tools Supporting AIRR Standards

Applications Supporting the Rearrangement Schema

The following list of software tools and databases support the TSV format of v1.2 of the AIRR Rearrangement schema.

Software	Version	Support	Reference
AIRR Python Li-	1.2	Input, output and validation	Vander Heiden et al. Front Im-
brary	1.2		munol, 2018. Vander Heiden et al. Front Im-
AIRR R Library	1.2	Input, output and validation	vander Heiden et al. Front Im- munol, 2018.
Alakazam	1.0.1	Input and output	Gupta & Vander Heiden et al.
			Bioinformatics, 2015.
Cell Ranger	4.0	Output	10x Genomics, Inc. Pleasanton,
			CA USA.
Change-O	0.4.2	Input, output and conversion	Gupta & Vander Heiden et al.
			Bioinformatics, 2015.
Decombinator	4.0.1	Output	Oakes et al. Front Immunol,
			2017.
IMGT/HighV-	1.7.0	Output	Alamyar et al. Methods Mol
QUEST			Biol, 2012.
IMGT/V-QUEST	3.5.16	Output	Giudicelli et al. Cold Spring
			Harb Protoc, 2011.
IgBLAST	1.11	Output	Ye et al. Nucleic Acids Res,
	0.6.7		2013.
immunarch	0.6.5	Input	ImmunoMind Team. 2019
ImmuneDB	0.24.0	Output	Rosenfeld et al. Front Immunol,
	0.07		2018.
immuneSIM	0.8.7	Output	Weber et al. Bioinformatics,
'D	2.0		2020.
iReceptor	3.0	Input and output	Corrie et al. Immunol Rev, 2018.
MiXCR	3.0.14	Outrast	Bolotin et al. Nat Methods,
MIACK	5.0.14	Output	2015.
RAbHIT	0.1.5	Input and output	Gidoni et al. Nat Commun,
KAUIIII	0.1.5	input and output	2019.
scirpy	0.3	Input	Sturm et al. Bioinformatics,
senpy	0.5	mpat	2020.
SCOPer	1.0.1	Input and output	Nouri & Kleinstein. Bioinfor-
	11011	input and curput	matics, 2018.
SHazaM	1.0.0	Input and output	Gupta & Vander Heiden et al.
		1 1	Bioinformatics, 2015.
SONAR	3	Output	Schramm et al. Front Immunol,
			2016.
sumrep	1.0	Input	Olson et al. Front Immunol,
			2019.
TIgGER	1.0.0	Input and output	Gadala-Maria et al. PNAS,
			2015.
TRIgS	2	Input	Lees & Shepherd. J Immunol
			Res, 2015.
VDJServer	1.2.0	Input and output	Christley et al. Front Immunol,
			2018
Vidjil-algo	2018.1	Output	Giraud et al. BMC Genomics,
X 71 4114 XX 7 4 354 0			2014.
Vidjil Web Platform	TBD	Input and conversion	Duez et al. PLoS ONE, 2016.
IGoR	TBD	Input and output	Marcou et al. Nat Commun,
OL CA			2018.
OLGA	TBD	Input and output	Sethna et al. Bioinformatics,
Dontio		Output	2019.
Partis	TBD	Output	Ralph & Matsen. PLoS Comput Biol, 2016. 14
2.5. Software			Biol, 2016. 149

AIRR Data Commons Repositories

These data repositories all implement the AIRR Data Commons (ADC) API programmatic access to query and down-load AIRR-seq data.

- iReceptor Public Archive
- VDJServer Community Data Portal

2.6 Community Resources

2.6.1 Resources and Tools Supporting AIRR Standards

2.6.2 Useful Websites for the AIRR Community

- The Antibody Society
- The AIRR Community of the Antibody Society
- B-T.CR Forum
- The AIRR Community GitHub
- The AIRR Standards GitHub Repository
- The AIRR Community Docker Hub

2.7 Appendix A: Key Terms

The following table provides definitions for terms and acronyms relevant to this documentation.

Term	Definition		
ADC	AIRR Data Commons		
AIRR	Adaptive Immune Receptor Repertoire		
AIRR-C	AIRR Community		
API	Application Programming Interface		
CAIRR	CEDAR AIRR		
CEDAR	Center for Expanded Data Annotation and Retrieval		
HTTP	Hypertext Transfer Protocol		
JSON	JavaScript Object Notation		
MiAIRR	Minimal Information about an Adaptive Immune Receptor Repertoire study		
REST	Representational State Transfer		
TSV	Tab Separated Values		
URL	Universal Resource Locator		
YAML	YAML Ain't Markup Language		

2.8 References

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[INSDC_FT] The DDBJ/ENA/GenBank Feature Table Definition. http://www.insdc.org/documents/feature-table

- [ENA_MANUAL] European Nucleotide Archive Annotated/Assembled Sequences User Manual. <http://ftp.ebi.ac. uk/pub/databases/ena/sequence/release/doc/usrman.txt>
- [GENBANK_FF] GenBank Flat File Format. <https://ftp.ncbi.nih.gov/genbank/gbrel.txt>
- [GENBANK_SR] GenBank Sample Record. < https://www.ncbi.nlm.nih.gov/Sitemap/samplerecord.html>
- [INSDC_XREF] Controlled vocabulary for /db_xref qualifier. ">http://www.insdc.org/documents/db_xref
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