

# Package: airr (via r-universe)

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**Type** Package

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**Title** AIRR Data Representation Reference Library

**Description** Schema definitions and read, write and validation tools  
for data formatted in accordance with the AIRR Data  
Representation schemas defined by the AIRR Community  
<<http://docs.airr-community.org>>.

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**URL** <http://docs.airr-community.org>

**BugReports** <https://github.com/airr-community/airr-standards/issues>

**BuildVignettes** true

**VignetteBuilder** knitr

**Encoding** UTF-8

**Depends** R (>= 3.1.2)

**Imports** jsonlite, methods, readr, stats, stringi, tools, yaml

**Suggests** knitr, rmarkdown, tibble, testthat

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**NeedsCompilation** no

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ExampleData	<i>Example AIRR data</i>
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## Description

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

## Format

- extdata/rearrangement-example.tsv.gz: Rearrangement TSV file.
- extdata/repertoire-example.yaml: Repertoire YAML file.
- extdata/germline-example.json: GermlineSet and GenotypeSet JSON file.

## Examples

```
# Load Rearrangement example
file <- system.file("extdata", "rearrangement-example.tsv.gz", package="airr")
rearrangement <- read_rearrangement(file)

# Load Repertoire example
file <- system.file("extdata", "repertoire-example.yaml", package="airr")
repertoire <- read_airr(file)

# Load GermlineSet and GenotypeSet examples
file <- system.file("extdata", "germline-example.json", package="airr")
germline <- read_airr(file)
```

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load_schema	<i>Load a schema definition</i>
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**Description**

load\_schema loads an AIRR object definition from the internal definition set.

**Usage**

```
load_schema(definition)
```

**Arguments**

definition      name of the schema definition.

**Details**

Valid definitions include:

- "Rearrangement"
- "Alignment"
- "Repertoire"
- "Study"
- "Subject"
- "Diagnosis"
- "Sample"
- "SampleProcessing"
- "DataProcessing"
- "GermlineSet"
- "GenotypeSet"

**Value**

A [Schema](#) object for the definition.

**See Also**

See [Schema](#) for the return object.

**Examples**

```
# Load the Rearrangement definition
schema <- load_schema("Rearrangement")

# Load the Repertoire definition
schema <- load_schema("Repertoire")
```

---

`read_airr`*Read an AIRR Data Model file in YAML or JSON format*

---

## Description

`read_airr` loads a YAML or JSON file containing AIRR Data Model records.

## Usage

```
read_airr(  
  file,  
  format = c("auto", "yaml", "json"),  
  validate = TRUE,  
  model = TRUE  
)
```

## Arguments

<code>file</code>	path to the input file.
<code>format</code>	format of the input file. Must be one of "auto", "yaml", or "json". If "auto" (default), the format will be detected from the file extension.
<code>validate</code>	run schema validation if TRUE.
<code>model</code>	if TRUE validate only AIRR DataFile defined objects. If FALSE attempt validation of all objects in data. Ignored if <code>validate=FALSE</code>

## Value

A named nested list contained in the AIRR Data Model with the top-level names reflecting the individual AIRR objects.

## See Also

See [Schema](#) for the AIRR schema definition objects. See [write\\_airr](#) for writing AIRR Data Model records in YAML or JSON format.

## Examples

```
# Get path to the Repertoire and GermlineSet example files  
f1 <- system.file("extdata", "repertoire-example.yaml", package="airr")  
f2 <- system.file("extdata", "germline-example.json", package="airr")  
  
# Load data files  
repertoire <- read_airr(f1)  
germline <- read_airr(f2)
```

---

read_tabular	<i>Read AIRR tabular data</i>
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### Description

read\_tabular reads a tab-delimited (TSV) file containing tabular AIRR records.

### Usage

```
read_tabular(file, schema, base = c("1", "0"), aux_types = NULL, ...)
```

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

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

### Arguments

file	input file path.
schema	Schema object defining the output format.
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).
aux_types	named vector or list giving the type for fields that are not defined in schema. The field name is the name, the value the type, denoted by one of "c" (character), "l" (logical), "i" (integer), "d" (double), or "n" (numeric).
...	additional arguments to pass to <a href="#">read_delim</a> .

### Details

read\_rearrangement reads an AIRR TSV containing Rearrangement data.

read\_alignment reads an AIRR TSV containing Alignment data.

### Value

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

### See Also

See [Schema](#) for the AIRR schema object definition. See [write\\_tabular](#) for writing AIRR 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)
```

---

Schema-class

*S4 class defining an AIRR standard schema*

---

**Description**

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

**Usage**

```
## S4 method for signature 'Schema'
names(x)

## S4 method for signature 'Schema,character'
x[i]

## S4 method for signature 'Schema'
x$name

InfoSchema

DataFileSchema

AlignmentSchema

RearrangementSchema

RepertoireSchema

GermlineSetSchema

GenotypeSetSchema

AIRRSchema
```

**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.

An object of class Schema of length 1.

An object of class Schema of length 1.

An object of class Schema of length 1.

An object of class Schema of length 1.

An object of class Schema of length 1.

An object of class list of length 26.

**Details**

The following predefined Schema objects are defined:

InfoSchema: AIRR Info Schema.

DataFileSchema: AIRR DataFile Schema.

AlignmentSchema: AIRR Alignment Schema.

RearrangementSchema: AIRR Rearrangement Schema.

RepertoireSchema: AIRR Repertoire Schema.

GermlineSetSchema: AIRR GermlineSet Schema.

GenotypeSetSchema: AIRR GenotypeSet Schema.

AIRRSchema: named list containing all non-experimental AIRR Schema objects.

**Slots**

definition name of the schema definition.

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.

---

validate_airr	<i>Validate an AIRR Data Model nested list representation</i>
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### Description

validate\_airr validates the fields in a named nested list representation of the AIRR Data Model. Typically, generating by reading of JSON or YAML formatted AIRR files.

### Usage

```
validate_airr(data, model = TRUE, each = FALSE)
```

### Arguments

data	list containing records of an AIRR Data Model object imported from a YAML or JSON representation.
model	if TRUE validate only AIRR DataFile defined objects. If FALSE attempt validation of all objects in data.
each	if TRUE return a logical vector with results for each object in data instead of a single TRUE or FALSE value.

### Value

Returns TRUE if the input data is compliant with AIRR standards and FALSE if not. If each=TRUE is set, then a vector with results for each each object in data is returned instead.

### See Also

See [Schema](#) for the AIRR schema definitions. See [read\\_airr](#) for loading AIRR Data Models from a file. See [write\\_airr](#) for writing AIRR Data Models to a file.

### Examples

```
# Get path to the rearrangement-example file
f1 <- system.file("extdata", "repertoire-example.yaml", package="airr")
f2 <- system.file("extdata", "germline-example.json", package="airr")

# Load data file
repertoire <- read_airr(f1)
germline <- read_airr(f2)

# Validate a single record
validate_airr(repertoire)

# Return validation for individual objects
validate_airr(germline, each=TRUE)
```



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validate_tabular	<i>Validate tabular AIRR data</i>
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### Description

validate\_tabular validates compliance of the contents of a data.frame to the AIRR standards.

### Usage

```
validate_tabular(data, schema)
```

```
validate_rearrangement(data)
```

### Arguments

data	data.frame of tabular data to validate.
schema	Schema object defining the data standard of the table.

### Details

validate\_rearrangement validates the standards compliance of AIRR Rearrangement data stored in a data.frame

### 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)

# Validate a data.frame against the Rearrangement schema
validate_rearrangement(df)
```

---

`write_airr`*Write AIRR Data Model records to YAML or JSON files*

---

### Description

`write_airr` writes a YAML or JSON file containing AIRR Data Model records.

### Usage

```
write_airr(  
  data,  
  file,  
  format = c("auto", "yaml", "json"),  
  validate = TRUE,  
  model = TRUE  
)
```

### Arguments

<code>data</code>	list containing AIRR Model Records.
<code>file</code>	output file name.
<code>format</code>	format of the output file. Must be one of "auto", "yaml", or "json". If "auto" (default), the format will be detected from the file extension.
<code>validate</code>	run schema validation prior to write if TRUE.
<code>model</code>	if TRUE validate and write only AIRR DataFile defined objects. If FALSE attempt validation and write of all objects in data.

### See Also

See [Schema](#) for the AIRR schema definition objects. See [read\\_airr](#) for reading to AIRR Data Model files.

### Examples

```
# Get path to the repertoire-example file  
file <- system.file("extdata", "repertoire-example.yaml", package="airr")  
  
# Load data file  
repertoire <- read_airr(file)  
  
# Write a Rearrangement data file  
outfile <- file.path(tempdir(), "output.yaml")  
write_airr(repertoire, outfile)
```

---

write_tabular	<i>Write an AIRR tabular data</i>
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---

### Description

write\_tabular writes a TSV containing AIRR tabular records.

### Usage

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

```
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.
schema	Schema object defining the output format.
base	starting index for positional fields in the output file. Fields in the input data are assumed to be 1-based closed-intervals (R style). If "1", then these fields will not be modified. If "0", then fields ending in <code>_start</code> and <code>_end</code> will be converted to 0-based half-open intervals (python style) in the output file.
...	additional arguments to pass to <a href="#">write_delim</a> .

### 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.

### See Also

See [Schema](#) for the AIRR schema object definition. See [read\\_tabular](#) for reading to AIRR files.

### 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_tabular(df, outfile, schema=RearrangementSchema)
```

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