## Package 'BiocIO'

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Title Standard Input and Output for Bioconductor Packages

**Version** 1.15.0

Description The `BiocIO` package contains high-level abstract classes and generics used by developers to build IO funcionality within the Bioconductor suite of packages. Implements `import()` and `export()` standard generics for importing and exporting biological data formats. `import()` supports whole-file as well as chunk-wise iterative import. The `import()` interface optionally provides a standard mechanism for 'lazy' access via `filter()` (on row or element-like components of the file resource), `select()` (on column-like components of the file resource) and `collect()`. The `import()` interface optionally provides transparent access to remote (e.g. via https) as well as local access. Developers can register a file extension, e.g., `.loom` for dispatch from character-based URIs to specific `import()` / `export()` methods based on classes representing file types, e.g., `LoomFile()`.

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Imports BiocGenerics, S4Vectors, methods, tools

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BiocFile-class BiocFile class objects

#### Description

BiocFile is the base virtual class for high-level file abstractions where subclasses are associated with a particular file format or type. It wraps a low-level representation of a file, currently either a path, URL, or connection object. We can represent a list of BiocFile objects with a BiocFileList.

#### Usage

```
BiocFileList(files)
resource(x)
resource(x) <- value
## S4 method for signature 'BiocFile'
resource(x)
## S4 replacement method for signature 'BiocFile,character_OR_connection'
resource(x) <- value
fileFormat(x)
## S4 method for signature 'character'
fileFormat(x)</pre>
```

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#### **BiocFile-class**

```
## S4 method for signature 'BiocFile'
fileFormat(x)
## S4 method for signature 'BiocFile'
path(object, ...)
## S4 method for signature 'BiocFile'
show(object)
FileForFormat(path, format = file_ext(path))
## S4 method for signature 'BiocFile'
as.character(x)
```

#### Arguments

files	character() A vector of file paths for the BiocFileList constructor
х	A BiocFile instance
object	A BiocFile instance
	additional arguments to lower-level functions, not used.
path, value	Either a character or connection object to replace the original resource
format	character(1) The file extension conducive to a file class name, e.g., CSVFile

#### Value

For constructors, an instance of that class. For extractors such as resource and path, typically a character vector of the file path. For FileForFormat, a convenient instance of the class for which the input file corresponds to.

#### **Accessor Methods**

In the code snippets below, x represents a BiocFile object.

- resource(x) Gets the low-level resource, either a character vector (a path or URL) or a connection.
- fileFormat(x) Gets a string identifying the file format. Can also be called directly on a character file path, in which case it uses a heuristic based on the file extension.

#### Author(s)

Michael Lawrence

#### See Also

Implementing classes include: BigWigFile, TwoBitFile, BEDFile, GFFFile, WIGFile

#### Examples

```
## For our examples, we create a class called CSVFILE that extends BiocFile
.CSVFile <- setClass("CSVFile", contains = "BiocFile")</pre>
## Constructor
CSVFile <- function(resource) {</pre>
    .CSVFile(resource = resource)
}
setMethod("import", "CSVFile", function(con, format, text, ...) {
    read.csv(resource(con), ...)
})
## Define export
setMethod("export", c("data.frame", "CSVFile"),
    function(object, con, format, ...) {
        write.csv(object, resource(con), ...)
    }
)
## Recommend CSVFile class for .csv files
temp <- tempfile(fileext = ".csv")</pre>
FileForFormat(temp)
## Create CSVFile
csv <- CSVFile(temp)</pre>
## Display path of file
path(csv)
## Display resource of file
resource(csv)
```

compression

File compression

#### Description

Methods and generics for file compression strategies.

#### Usage

```
decompress(manager, con, ...)
## S4 method for signature 'ANY'
decompress(manager, con, ...)
## S4 method for signature 'CompressedFile'
```

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```
decompress(manager, con, ...)
## S4 method for signature 'character'
decompress(manager, con, ...)
## S4 method for signature 'CompressedFile'
fileFormat(x)
```

#### Arguments

manager	The connection manager, defaults to the internal manager class
con	The connection from which data is loaded or to which data is saved. If this is a character vector, it is assumed to be a file name and a corresponding file connection is created and then closed after exporting the object. If it is a BiocFile derivative, the data is loaded from or saved to the underlying resource. If missing, the function will return the output as a character vector, rather than writing to a connection.
	Parameters to pass to the format-specific method.
x	A BiocFile instance

#### Value

A decompressed representation of a CompressedFile or character object

#### **Related functions**

#### Examples

```
file <- tempfile(fileext = ".gzip")
decompress(con = file)</pre>
```

I0

Import and export

#### Description

The functions import and export load and save objects from and to particular file formats.

#### Usage

import(con, format, text, ...) ## S4 method for signature 'connection, character, ANY' import(con, format, text, ...) ## S4 method for signature 'connection,missing,ANY' import(con, format, text, ...) ## S4 method for signature 'character,missing,ANY' import(con, format, text, ...) ## S4 method for signature 'character, character, ANY' import(con, format, text, ...) ## S4 method for signature 'missing, ANY, character' import(con, format, text, ...) export(object, con, format, ...) ## S4 method for signature 'ANY, connection, character' export(object, con, format, ...) ## S4 method for signature 'ANY,connection,missing' export(object, con, format, ...) ## S4 method for signature 'ANY,missing,character' export(object, con, format, ...) ## S4 method for signature 'ANY, character, missing' export(object, con, format, ...) ## S4 method for signature 'ANY, character, character' export(object, con, format, ...) ## S4 method for signature 'CompressedFile,missing,ANY' import(con, format, text, ...) ## S4 method for signature 'ANY,CompressedFile,missing' export(object, con, format, ...)

#### Arguments

con

The connection from which data is loaded or to which data is saved. If this is a character vector, it is assumed to be a file name and a corresponding file connection is created and then closed after exporting the object. If it is a BiocFile derivative, the data is loaded from or saved to the underlying resource. If missing, the function will return the output as a character vector, rather than

	writing to a connection.
format	The format of the output. If missing and con is a file name, the format is derived from the file extension. This argument is unnecessary when con is a derivative of BiocFile.
text	If con is missing, this can be a character vector directly providing the string data to import.
	Parameters to pass to the format-specific method.
object	The object to export.

writing to a connection

#### Value

If con is missing, a character vector containing the string output. Otherwise, nothing is returned.

#### Author(s)

Michael Lawrence

#### See Also

Format-specific options for the popular formats: GFF, BED, BED15, BEDGRAPH, WIG, BIGWIG

#### Examples

```
## To illustrate export(), import(), and yeild(), we create a class, CSVFILE
.CSVFile <- setClass("CSVFile", contains = "BiocFile")</pre>
## Constructor
CSVFile <- function(resource) {</pre>
    .CSVFile(resource = resource)
}
## Define import
setMethod("import", "CSVFile",
    function(con, format, text, ...) {
        read.csv(resource(con), ...)
    }
)
## Define export
setMethod("export", c("data.frame", "CSVFile"),
    function(object, con, format, ...) {
        write.csv(object, resource(con), ...)
    }
)
## Usage
temp <- tempfile(fileext = ".csv")</pre>
csv <- CSVFile(temp)</pre>
export(mtcars, csv)
```

df <- import(csv)</pre>

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