

# Package ‘alabaster.files’

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**Title** Wrappers to Save Common File Formats

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**Description** Save common bioinformatics file formats within the alabaster framework.  
This includes BAM, BED, VCF, bigWig, bigBed, FASTQ, FASTA and so on.  
We save and load additional metadata for each file,  
and we support linkage between each file and its corresponding index.

**Depends** alabaster.base,

**Imports** methods, S4Vectors, BiocGenerics, Rsamtools

**Suggests** BiocStyle, rmarkdown, knitr, testthat, VariantAnnotation,  
rtracklayer, Biostrings

**VignetteBuilder** knitr

**RoxygenNote** 7.2.3

**biocViews** DataRepresentation, DataImport

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BamFileReference	<i>Reference to a BAM file</i>
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## Description

Reference to a BAM file, for saving and reading in the **alabaster** framework.

## Usage

```
BamFileReference(path, index = NULL)
```

## Arguments

path	String containing the path to a BAM file.
index	String specifying the path to a BAI or CSI index file, or NULL if no index is available.

## Value

A BamFileReference instance that can be used in `saveObject`.

## Author(s)

Aaron Lun

## Examples

```
# Using a BAM file from Rsamtools.
fl <- system.file("extdata", "ex1.bam", package="Rsamtools", mustWork=TRUE)

# Creating a BamFileReference.
wrapped <- BamFileReference(fl)
wrapped

# Fetching the path information:
path(wrapped)
wrapped$index
```

```
# Saving to disk:
dir <- tempfile()
saveObject(wrapped, dir)
list.files(dir, recursive=TRUE)

# Reading it back again:
readObject(dir)
```

---

BcfFileReference	<i>Reference to a BCF file</i>
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## Description

Reference to a BCF file, for saving and loading in the **alabaster** framework.

## Usage

```
BcfFileReference(path, index = NULL)
```

## Arguments

path	String containing the path to a BCF file.
index	String specifying the path to an index file in tabix or CSI format, or NULL if no index is available.

## Value

A BcfFileReference instance that can be used in [saveObject](#).

## Author(s)

Aaron Lun

## Examples

```
# Using Rsamtools's example file.
fl <- system.file("extdata", "ex1.bcf.gz", package="Rsamtools")

# Creating a BcfFileReference.
wrapped <- BcfFileReference(fl)
wrapped

# Fetching the path information:
path(wrapped)
wrapped$index

# Staging the BcfFileReference.
```

```
dir <- tempfile()
saveObject(wrapped, dir)
list.files(dir, recursive=TRUE)

# Loading it back again:
readObject(dir)
```

---

BedFileReference	<i>Reference to a BED file</i>
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---

### Description

Reference to a BED file, for saving and loading in the **alabaster** framework.

### Usage

```
BedFileReference(path, index = NULL)
```

### Arguments

path	String containing the path to a Gzip- or BGZF-compressed BED file.
index	String containing a path to a tabix file. If supplied, path should be coordinate-sorted and BGZF-compressed.

### Value

A BedFileReference instance that can be used in [saveObject](#).

### Author(s)

Aaron Lun

### Examples

```
# Mocking up a BED file.
raw <- tempfile(fileext=".bed")
bed <- write("chr1\t2222\t33333", file=raw)
tmp <- tempfile(fileext=".bed.bgz")
Rsamtools::bgzip(raw, tmp)

# Creating a BedFileReference.
wrapped <- BedFileReference(tmp)
wrapped

# Extracting the paths:
path(wrapped)
wrapped$index
```

```
# Saving it to disk.
dir <- tempfile()
saveObject(wrapped, dir)
list.files(dir, recursive=TRUE)

# Loading it back again:
readObject(dir)
```

---

BgzipIndexWrapper      *Wrapper for a Bgzip index file*

---

### Description

This class is deprecated and only listed here for back-compatibility purposes.

### Usage

```
BgzipIndexWrapper(path)
```

### Arguments

path                      String containing the path to a Bgzip index file.

### Details

The BgzipIndexWrapper class is a subclass of a [Wrapper](#), so all of the methods of the latter can also be used here, e.g., path.

### Value

A BgzipIndexWrapper instance that can be used in [stageObject](#).

### Author(s)

Aaron Lun

### Examples

```
# Mocking up a FASTA index file.
input <- system.file("extdata", "ce2dict1.fa", package="Rsamtools")
temp <- tempfile(fileext=".fa.bgz")
copy <- Rsamtools::bgzip(input, dest=temp)
Rsamtools::indexFa(copy)

# Creating a BgzipIndexWrapper.
wrapped <- BgzipIndexWrapper(paste0(copy, ".gzi"))
wrapped

# Staging the BgzipIndexWrapper.
```

```
dir <- tempfile()
library(alabaster.base)
info <- stageObject(wrapped, dir, "tab")
invisible(.writeMetadata(info, dir))
list.files(dir, recursive=TRUE)

# Loading it back again:
meta <- acquireMetadata(dir, "tab/file.fa.bgz.gzi")
loadObject(meta, dir)
```

---

BigBedFileReference    *Reference to a bigBed file*

---

### Description

Reference to a bigBed file, for saving and loading in the **alabaster** framework.

### Usage

```
BigBedFileReference(path)
```

### Arguments

path                    String containing the path to a bigBed file.

### Value

A BigBedFileReference instance that can be used in [stageObject](#).

### Author(s)

Aaron Lun

### Examples

```
# Mocking up a bigBed file.
test_path <- system.file("tests", "test.bb", package = "rtracklayer")

# Creating a BigBedFileReference.
wrapped <- BigBedFileReference(test_path)
wrapped

# Staging the BigBedFileReference.
dir <- tempfile()
saveObject(wrapped, dir)
list.files(dir, recursive=TRUE)

# Loading it back again:
readObject(dir)
```

---

BigWigFileReference     *Reference to a bigWig file*

---

## Description

Reference to a bigWig file, for saving and loading in the **alabaster** framework.

## Usage

```
BigWigFileReference(path)
```

## Arguments

path                    String containing the path to a bigWig file.

## Value

A BigWigFileReference instance that can be used in [stageObject](#).

## Author(s)

Aaron Lun

## Examples

```
# Mocking up a bigWig file.
test_path <- system.file("tests", "test.bw", package = "rtracklayer")

# Creating a BigWigFileReference.
wrapped <- BigWigFileReference(test_path)
wrapped

# Staging the BigWigFileReference.
dir <- tempfile()
saveObject(wrapped, dir)
list.files(dir, recursive=TRUE)

# Loading it back again:
readObject(dir)
```

---

FastaFileReference      *Reference to a FASTA file*

---

### Description

Reference to a FASTA file, for saving and loading in the **alabaster** framework.

### Usage

```
FastaFileReference(path, seqtype = "DNA", faindex = NULL, gzindex = NULL)
```

### Arguments

path	String containing the path to a Gzip- or BGZF-compressed FASTA file.
seqtype	String specifying the sequence type. This should be one of "DNA", "RNA", "AA" or "custom".
faindex	String specifying the path to an FASTA index file, or NULL if no index is available. If an index is supplied, the file at path should be BGZF-compressed, and gzindex should also be supplied.
gzindex	String specifying the path to a BGZF index file, or NULL if no index is available. If an index is supplied, the file at path should be BGZF-compressed, and faindex should also be supplied.

### Value

A FastaFileReference instance that can be used in [saveObject](#).

### Author(s)

Aaron Lun

### Examples

```
# Mocking up a FASTA file.
tmp <- tempfile(fileext=".fa.gz")
write(">FOOBAR\nacgtacgt", gzfile(tmp))

# Creating a FastaFileReference.
wrapped <- FastaFileReference(tmp)
wrapped

# Saving to disk:
dir <- tempfile()
saveObject(wrapped, dir)
list.files(dir, recursive=TRUE)

# Loading it back again:
readObject(dir)
```



---

FastqFileReference      *Reference to a FASTQ file*

---

## Description

Reference to a FASTQ file, for saving and loading in the **alabaster** framework.

## Usage

```
FastqFileReference(  
    path,  
    seqtype = "DNA",  
    qualtype = "phred",  
    qualoffset = 33,  
    faindex = NULL,  
    gzindex = NULL  
)
```

## Arguments

path	String containing the path to a Gzip- or BGZF-compressed FASTQ file.
seqtype	String specifying the sequence type. This should be one of "DNA", "RNA", "AA" or "custom".
qualtype	String specifying the type of the quality strings. This should be one of "phred" or "solexa".
qualoffset	Integer specifying the encoding offset for the quality strings. This is only used when qualtype="phred", in which case it should either be 33 or 64.
faindex	String specifying the path to an FASTA index file, or NULL if no index is available. If an index is supplied, the file at path should be BGZF-compressed, and gzindex should also be supplied.
gzindex	String specifying the path to a BGZF index file, or NULL if no index is available. If an index is supplied, the file at path should be BGZF-compressed, and faindex should also be supplied.

## Value

A FastqFileReference instance that can be used in [saveObject](#).

## Author(s)

Aaron Lun

**Examples**

```
# Mocking up a FASTQ file.
tmp <- tempfile(fileext=".fq.gz")
write("@FOOBAR\nacgtacgt\n+134987382", gzfile(tmp))

# Creating a FastqFileReference.
wrapped <- FastqFileReference(tmp)
wrapped

# Staging the FastqFileReference.
dir <- tempfile()
saveObject(wrapped, dir)
list.files(dir, recursive=TRUE)

# Loading it back again:
readObject(dir)
```

---

FileReference	<i>Virtual file reference class</i>
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**Description**

A virtual class for file reference objects. This implements common methods for `path`, `[[` and `$`.

**Author(s)**

Aaron Lun

---

GffFileReference	<i>Reference to a GFF file</i>
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---

**Description**

Reference to a GFF2/3 file, for saving and loading in the **alabaster** framework.

**Usage**

```
GffFileReference(path, index = NULL)
```

**Arguments**

<code>path</code>	String containing the path to a Gzip- or BGZF-compressed GFF file. The format is automatically detected from the file extension ( <code>.gff2</code> , <code>.gff3</code> or <code>.gtf</code> ).
<code>index</code>	String specifying the path to a tabix file in tabix format, or NULL if no index is available. If supplied, <code>path</code> should be coordinate-sorted and BGZF-compressed.

**Value**

A GffFileReference instance that can be used in [saveObject](#).

**Author(s)**

Aaron Lun

**Examples**

```
# Using rtracklayer's example GFF file.
src <- system.file("tests", "genes.gff3", package = "rtracklayer")
fl <- tempfile(fileext=".gff3.gz")
writeLines(con=gzfile(fl), readLines(src))

# Creating a GffFileReference.
wrapped <- GffFileReference(fl)
wrapped

# Saving it:
dir <- tempfile()
saveObject(wrapped, dir)
list.files(dir, recursive=TRUE)

# Loading it back again:
readObject(dir)
```

---

GmtFileReference	<i>Reference to a GMT file</i>
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---

**Description**

Reference to a GMT file, for saving and loading in the **alabaster** framework.

**Usage**

```
GmtFileReference(path)
```

**Arguments**

path                   String containing the path to a Gzip-compressed GMT file.

**Value**

A GmtFileReference instance that can be used in [stageObject](#).

**Author(s)**

Aaron Lun

## Examples

```
# Mocking up a GMT file.
tmp <- tempfile(fileext=".gmt.gz")
write("SET1\tdescription\tgene1\tgene2\tgene3", file=gzfile(tmp))

# Creating a GmtFileReference.
wrapped <- GmtFileReference(tmp)
wrapped

# Saving to disk:
dir <- tempfile()
saveObject(wrapped, dir)

# Loading it back again:
readObject(dir)
```

---

TabixIndexWrapper	<i>Wrapper for a Tabix file</i>
-------------------	---------------------------------

---

## Description

This class is deprecated and only listed here for back-compatibility purposes.

## Usage

```
TabixIndexWrapper(path)
```

## Arguments

path	String containing the path to a Tabix file.
------	---

## Details

The `TabixIndexWrapper` class is a subclass of a [Wrapper](#), so all of the methods of the latter can also be used here, e.g., `path`.

## Value

A `TabixIndexWrapper` instance that can be used in `stageObject`.

## Author(s)

Aaron Lun

## Examples

```
# Mocking up a Tabix file.
test_tbx <- system.file("extdata", "example.gtf.gz.tbi", package="Rsamtools")

# Creating a TabixIndexWrapper.
wrapped <- TabixIndexWrapper(test_tbx)
wrapped

# Staging the TabixIndexWrapper.
dir <- tempfile()
library(alabaster.base)
info <- stageObject(wrapped, dir, "tab")
invisible(.writeMetadata(info, dir))
list.files(dir, recursive=TRUE)

# Loading it back again:
meta <- acquireMetadata(dir, "tab/file.tbi")
loadObject(meta, dir)
```

---

virtual-wrapper

*Virtual wrapper classes*

---

## Description

Defines some base classes for the concrete wrappers for specific file formats. This provides a standard set of methods that can be applied to all Wrapper instances.

## Wrapper methods

Any instance  $x$  of a base Wrapper class can be used with the `path(x)` method, which returns a string containing the path to the file on the current file system.

The Wrapper class inherits from the [Annotated](#) class, so users can also get and set metadata via `metadata(x)`.

## IndexedWrapper methods

The IndexedWrapper class inherits from the Wrapper class and can be used with all its methods. It additionally implements the `index(x)` method, which returns another Wrapper object for the associated index file (or NULL, if no index file exists).

## CompressedWrapper methods

The CompressedWrapper class inherits from the Wrapper class and can be used with all its methods. It additionally implements the `compression(x)` method, which returns a string specifying the compression strategy.

**CompressedIndexedWrapper methods**

The CompressedIndexedWrapper class inherits from both the IndexedWrapper and CompressedWrapper classes and can be used with all their methods.

**Author(s)**

Aaron Lun

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