Package 'alabaster.string'

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Title Save and Load Biostrings to/from File

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Description Save Biostrings objects to file artifacts, and load them back into memory. This is a more portable alternative to serialization of such objects into RDS files. Each artifact is associated with metadata for further interpretation; downstream applications can enrich this metadata with context-specific properties.

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Depends Biostrings, alabaster.base

Imports methods

Suggests BiocStyle, rmarkdown, knitr, testthat

VignetteBuilder knitr

RoxygenNote 7.2.1

biocViews DataImport, DataRepresentation

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loadXStringSet

Description

Load a DNAStringSet object, typically from a FASTA or FASTQ file generated by the corresponding stageObject method.

Usage

loadXStringSet(seq.info, project)

Arguments

seq.info	A named list of metadata for this object.
project	Any argument accepted by the acquisition functions, see ?acquireFile. By
	default, this should be a string containing the path to a staging directory.

Value

A DNAStringSet containing DNA sequences. This may also be a QualityScaledDNAStringSet with quality scores.

Examples

```
library(Biostrings)
stuff <- DNAStringSet(c("AAA", "CC", "G", "TTTT"))</pre>
```

```
tmp <- tempfile()
dir.create(tmp)
info <- stageObject(stuff, tmp, path="dna_thing")
loadXStringSet(info, project=tmp)</pre>
```

Description

Stage a XStringSet by saving it to the appropriate file format.

Usage

```
## S4 method for signature 'XStringSet'
stageObject(x, dir, path, child = FALSE, ...)
```

Arguments

х	A XStringSet or any of its subclasses, in particular a QualityScaledXStringSet.
dir	String containing the path to the staging directory.
path	String containing a prefix of the relative path inside dir where x is to be saved. The actual path used to save x may include additional components, see Details.
child	Logical scalar indicating whether x is a child of a larger object.
	Further arguments to pass to specific methods.

Value

A list containing metadata for x. A subdirectory is created at path inside dir and the contents of x are saved to various files within that subdirectory. If x is a QualityScaledXStringSet, a FASTQ file is created instead of a FASTA file.

Author(s)

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Examples

```
library(Biostrings)
stuff <- DNAStringSet(c("AAA", "CC", "G", "TTTT"))
tmp <- tempfile()
dir.create(tmp)
stageObject(stuff, tmp, path="dna_thing")
list.files(tmp, recursive=TRUE)</pre>
```

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