# Package 'alabaster.matrix'

September 24, 2024

Title Load and Save Artifacts from File

**Version** 1.5.10

Date 2024-09-22

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Description

Save matrices, arrays and similar objects into 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.

Depends alabaster.base

**Imports** methods, BiocGenerics, S4Vectors, DelayedArray (>= 0.31.8), S4Arrays, SparseArray (>= 1.5.22), rhdf5 (>= 2.47.1), HDF5Array, Matrix, Rcpp

Suggests testthat, knitr, BiocStyle, chihaya, BiocSingular, ResidualMatrix

LinkingTo Rcpp

VignetteBuilder knitr

RoxygenNote 7.3.2

**Encoding** UTF-8

biocViews DataImport, DataRepresentation

git\_url https://git.bioconductor.org/packages/alabaster.matrix

git\_branch devel

git\_last\_commit 5d826ee

git\_last\_commit\_date 2024-09-22

**Repository** Bioconductor 3.20

**Date/Publication** 2024-09-24

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AmalgamatedArray *Amalgamated array class* 

# Description

Implements an amalgamated array, equivalent to a delayed combination of DelayedArray objects. It allows stageObject to save a combination of multiple matrices without actually aggregating their data into a single file.

#### Constructors

AmalgamatedArraySeed(..., along=1) accepts any number of named array-like objects and returns a AmalgamatedArraySeed. Each object corresponds to a block and should be named accordingly; names should be unique and non-empty. The along argument specifies the dimension in which matrices should be combined - for matrices, this is 1 for rows, 2 for columns.

AmalgamatedArray(..., along=1) accepts any number of named array-like objects and returns a AmalgamatedArray. Alternatively, a single AmalgamatedArraySeed may be provided in ....

# Functions

componentNames(x) will return a character vector of names of component arrays in a AmalgamatedArray(Seed) object x.

extractComponents(x) will return a named list of array-like objects, corresponding to the component arrays used to construct the AmalgamatedArray(Seed) object x.

stageObject(x, dir, path, child = FALSE) will save the AmalgamatedArray x and its components into the path inside dir. Each component array is staged into its own subdirectory inside path.

#### **Comments on usage**

The AmalgamatedArraySeed is closely related to (and in fact, is a subclass of) the DelayedAbind class. This means that we can leverage many of the **DelayedArray** methods for handling the delayed bind. In theory, we could just use a DelayedAbind directly and save it with **chihaya** in stageObject (via preserveDelayedOperations(TRUE)). However, this provides fewer opportunities for tracking and manipulating the samples. It also saves the per-sample matrices into a single file, which eliminates possibilities for per-file deduplication and linking, e.g., with recycleHdf5Files(TRUE).

## createRawArraySeed

# Author(s)

Aaron Lun

# Examples

```
first <- Matrix::rsparsematrix(10, 10, 0.1)
second <- Matrix::rsparsematrix(10, 20, 0.1)
mat <- AmalgamatedArray(list(foo = first, bar = second), along=2)
mat
componentNames(mat)
out <- extractComponents(mat)
lapply(out, dim)</pre>
```

createRawArraySeed Array loading utilities

# Description

Utilities for loading an array saved by stageObject.

## Usage

```
.createRawArraySeed(info, project, names = TRUE)
```

```
.extractArrayDimnames(path, group, ndim)
```

# Arguments

info	A named list of metadata for this array.
project	Any argument accepted by the acquisition functions, see ?acquireFile. By default, this should be a string containing the path to a staging directory.
names	Logical scalar indicating whether the seed should be annotated with dimnames (if available).
path	String containing the path to the file containing said array.
group	String containing the name of the group with the dimnames.
ndim	Integer scalar specifying the number of dimensions.

# Details

For .createArraySeed, the array should be one of:

- hdf5\_dense\_array
- hdf5\_sparse\_matrix
- hdf5\_delayed\_array
- amalgamated\_array

For delayed arrays, the file may contain a seed array with the "custom alabaster local array" type. This should have a path dataset containing a relative path to another array in the same project, which is loaded and used as the seed for this delayed array. Callers can overwrite this behavior by setting "custom alabaster local array" in the knownArrays from **chihaya** before calling .createRawArraySeed.

For .extractArrayDimnames, path is expected to be a HDF5 file with a group specified by group. Each child of this group is a string dataset named after a (0-indexed) dimension, containing the names for that dimension.

## Value

.createRawArraySeed returns a seed that can be used in the DelayedArray constructor.

.extractArrayDimnames returns a list of character vectors or NULL, containing the dimnames.

#### Author(s)

Aaron Lun

# Examples

```
# Staging an array as an example:
dir <- tempfile()
dir.create(dir)
mat <- array(rpois(10000, 10), c(50, 20, 10))
meta <- stageObject(mat, dir, "whee")
# Loading it back as a DelayedArray seed:
.createRawArraySeed(meta, project=dir)
```

DelayedMask

Delayed masking

# Description

Delayed masking of missing values, based on replacement of placeholder values with NA. This allows missingness to be encoded in frameworks without the same concept of NA as R.

# Usage

```
DelayedMask(x, placeholder)
```

## Arguments

x	An existing <b>DelayedArray</b> seed.
placeholder	Placeholder value to replace with NA. This should be of the same type as $type(x)$ .

#### preserveDelayedOperations

#### Details

If is.na(placeholder) is true for double-precision x, masking is performed for all values of x where is.na is true. This includes both NaNs and NAs; no attempt is made to distinguish between the NaN payloads.

Currently, an error is raised for any integer x that produces non-missing values of  $-2^{31}$  without a placeholder of NA\_integer\_. This is because R cannot distinguish the integer  $-2^{31}$  from an integer-type NA.

# Value

A DelayedMask object, to be wrapped in a DelayedArray.

#### Author(s)

Aaron Lun

#### Examples

```
original <- DelayedArray(matrix(rpois(40, lambda=2), ncol=5))
original
masked <- DelayedMask(original, 0)
DelayedArray(masked)</pre>
```

preserveDelayedOperations

Preserve delayed operations during staging

# Description

Preserve delayed operations via chihaya when staging a DelayedArray with stageObject.

# Usage

preserveDelayedOperations(preserve)

#### Arguments

preserve Whether to preserve delayed operations using the **chihaya** specification.

# Details

By default, any DelayedArray in stageObject will be saved as a new dense array or sparse matrix. However, if this option is enabled, DelayedArrays will instead be saved in the **chihaya** specification, where the delayed operations are themselves stored in the HDF5 file (see https://artifactdb.github.io/chihaya/ for details).

The **chihaya** specification is more complicated to parse but can be helpful in reducing disk usage. One simple example is to avoid sparsity-breaking or integer-to-float operations by storing their delayed representations in the file. If the seed matrix is derived from some immutable reference location, advanced users can even store links to that location instead of duplicating the seed data.

# Value

Logical scalar indicating whether delayed operations are to be preserved by the DelayedArray method. If preserve is supplied, it is used to set this scalar, and the *previous* value of the scalar is invisibly returned.

# Author(s)

Aaron Lun

# Examples

```
preserveDelayedOperations()
old <- preserveDelayedOperations(TRUE)
preserveDelayedOperations()
preserveDelayedOperations(old)</pre>
```

readArray

Read a dense array from disk

## Description

Read a dense high-dimensional array from its on-disk representation. This is usually not directly called by users, but is instead called by dispatch in readObject.

# Usage

```
readArray(path, metadata, ...)
```

## Arguments

path	String containing a path to a directory, itself created by the saveObject method for a dense array.
metadata	Named list of metadata for this object, see readObject for more details.
	Further arguments, ignored.

## Value

A dense file-backed ReloadedArray.

#### Author(s)

Aaron Lun

# See Also

"saveObject, array-method", to create the directory and its contents.

#### readDelayedArray

## Examples

```
arr <- array(rpois(10000, 10), c(50, 20, 10))
dimnames(arr) <- list(
    paste0("GENE_", seq_len(nrow(arr))),
    letters[1:20],
    NULL
)
dir <- tempfile()
saveObject(arr, dir)
readObject(dir)</pre>
```

readDelayedArray Read a delayed array from disk

# Description

Read a delayed high-dimensional array from its on-disk representation. This is usually not directly called by users, but is instead called by dispatch in readObject.

# Usage

```
readDelayedArray(path, metadata, delayed_array.reload.args = list(), ...)
```

# Arguments

path	String containing a path to a directory, itself created by the saveObject method for a delayed array.
metadata	Named list of metadata for this object, see readObject for more details.
delayed_array.	reload.args Named list of arguments to be passed to reloadDelayedObject.
	Further arguments, ignored.

# Value

A multi-dimensional array-like object.

# Author(s)

Aaron Lun

# See Also

"saveObject,DelayedArray-method", to create the directory and its contents.

reloadDelayedObject, for the methods to reload each delayed operation.

# Examples

```
arr <- array(rpois(10000, 10), c(50, 20, 10))
dimnames(arr) <- list(
    paste0("GENE_", seq_len(nrow(arr))),
    letters[1:20],
    NULL
)
dir <- tempfile()
saveObject(arr, dir)
readObject(dir)</pre>
```

readSparseMatrix Read a sparse matrix from disk

## Description

Read a sparse matrix from its on-disk representation. This is usually not directly called by users, but is instead called by dispatch in readObject.

# Usage

```
readSparseMatrix(path, metadata, ...)
```

# Arguments

path	String containing a path to a directory, itself created by the saveObject method for a spars matrix.
metadata	Named list of metadata for this object, see readObject for more details.
	Further arguments, ignored.

# Value

A sparse ReloadedMatrix object.

# Author(s)

Aaron Lun

# See Also

"saveObject, sparseMatrix-method", to create the directory and its contents.

#### Examples

```
mat <- Matrix::rsparsematrix(100, 200, density=0.2)
rownames(mat) <- paste0("GENE_", seq_len(nrow(mat)))
dir <- tempfile()
saveObject(mat, dir)
readObject(dir)</pre>
```

recycleHdf5Files Recycle existing HDF5 files

# Description

Re-use existing files in HDF5-backed arrays rather than reserializing them in stageObject.

# Usage

```
recycleHdf5Files(recycle)
```

### Arguments

recycle Whether to recycle existing files for HDF5-backed DelayedArrays.

# Details

If this options is enabled, stageObject will attempt to link/copy existing files for any HDF5-backed DelayedArray instances - most specifically, HDF5Array objects and H5SparseMatrix objects using the 10X format. This avoids re-serialization of the data for faster staging. It also allows advanced users to add their own customizations into the HDF5 file during staging, as long as they do not interfere with loadArray.

By default, this option is disabled as the properties of the existing file are not known in the general case. In particular, the file might contain other groups/datasets that are irrelevant, and use up extra disk space if copied; or confidential, and should not be stored in the staging directory. Users should only enable this option if they have full control over the generation and contents of the backing HDF5 files.

Also note that any dimnames on x will be ignored during recycling.

# Value

Logical scalar indicating whether HDF5 files are to be reused. If recycle is supplied, it is used to set this scalar, and the *previous* value of the scalar is invisibly returned.

# Author(s)

Aaron Lun

```
recycleHdf5Files()
old <- recycleHdf5Files(TRUE)
recycleHdf5Files()
recycleHdf5Files(old)</pre>
```

ReloadedArraySeed Reloaded alabaster array

#### Description

An array that was reloaded from disk by the readObject function. This allows methods to refer to the existing on-disk representation by inspecting the path. For example, saveObject can just copy/link to the files instead of repeating the saving process.

#### Usage

```
ReloadedArraySeed(path, seed = NULL, ...)
```

```
ReloadedArray(path, seed = NULL, ...)
```

#### Arguments

path	String containing a path to the directory with the on-disk array representation. Alternatively an existing ReloadedArraySeed, which is returned without modification.
seed	Contents of the loaded array, e.g., as an ordinary R array, a DelayedArray or a sparse matrix. If NULL, this is obtained by calling readObject.
	Further arguments to pass to readObject when seed=NULL.

# Details

The ReloadedArraySeed is a subclass of the WrapperArraySeed and will just forward all operations to the underlying seed. Its main purpose is to track the path that was originally used to generate seed, which enables optimizations for methods that need to operate on the files.

One obvious optimization is the specialization of saveObject on ReloadedArray instances. Instead of loading the array data back into the R session and saving it again, the saveObject method can just link or copy the existing files. This behavior is controlled by the optional ReloadedArray.reuse.files option in the saveObject method, which can be one of:

- "copy": copy the files from the original directory (as stored in the ReloadedArray object) to the new path specified in saveObject.
- "link": create a hard link from the files in the original directory to the new path. If this fails, we silently fall back to a copy. This mode is the default approach.
- "symlink": create a symbolic link from the files in the original directory to the new path. Each symbolic link refers to an absolute path in the original directory, which is useful when the contents of path might be moved (but the original directory will not).
- "relsymlink": create a symbolic link from the files in the original directory to the new path. Each symbolic link refers to an relative path to its corresponding file in the original directory, which is useful when both path and the original directory are moved together, e.g., as they are part of the same parent object like a SummarizedExperiment.
- "none": ignore existing files and just save the contents by calling "saveObject, DelayedArray-method".

## Value

For the constructors, an instance of the ReloadedArraySeed or ReloadedArray.

#### saveArray

#### Examples

```
arr <- array(rpois(10000, 10), c(50, 20, 10))
dir <- tempfile()
saveObject(arr, dir)
obj <- readArray(dir)
obj
DelayedArray::showtree(obj)</pre>
```

saveArray

Save a multi-dimensional array to disk

# Description

Save a high-dimensional array to its on-disk representations.

# Usage

```
## S4 method for signature 'array'
saveObject(x, path, ...)
## S4 method for signature 'denseMatrix'
saveObject(x, path, ...)
```

### Arguments

Х	An integer, numeric, logical or character array. Alternatively, any of the dense-
	Matrix subclasses from the Matrix package.
path	String containing the path to a directory in which to save x.
	Further arguments, currently ignored.

# Value

x is saved to path and NULL is invisibly returned.

# Author(s)

Aaron Lun

## See Also

readArray, to read the directory contents back into the R session.

```
mat <- array(rpois(10000, 10), c(50, 20, 10))
dimnames(mat) <- list(
    paste0("GENE_", seq_len(nrow(mat))),
    letters[1:20],
    NULL
)</pre>
```

```
dir <- tempfile()
saveObject(mat, dir)
list.files(dir)</pre>
```

saveDelayedArray Save DelayedArrays to disk

# Description

Save DelayedArray objects to their on-disk representation.

# Usage

```
## S4 method for signature 'DelayedArray'
saveObject(
    x,
    path,
    DelayedArray.dispatch.pristine = TRUE,
    DelayedArray.preserve.ops = FALSE,
    DelayedArray.store.args = list(),
    ...
)
```

# Arguments

pristine arrays. DelayedArray.preserve.ops	Х	A DelayedArray object.			
Logical scalar indicating whether to call the saveObject methods of seeds of pristine arrays. DelayedArray.preserve.ops Logical scalar indicating whether delayed operations should be preserved on disk. DelayedArray.store.args Named arguments to pass to storeDelayedObject.	path	String containing a path to a directory in which to save x.			
pristine arrays. DelayedArray.preserve.ops Logical scalar indicating whether delayed operations should be preserved on disk. DelayedArray.store.args Named arguments to pass to storeDelayedObject.	DelayedArray.di	spatch.pristine			
Logical scalar indicating whether delayed operations should be preserved on disk. DelayedArray.store.args Named arguments to pass to storeDelayedObject.		Logical scalar indicating whether to call the saveObject methods of seeds of pristine arrays.			
disk. DelayedArray.store.args Named arguments to pass to storeDelayedObject.	DelayedArray.preserve.ops				
Named arguments to pass to storeDelayedObject.		Logical scalar indicating whether delayed operations should be preserved on- disk.			
	DelayedArray.store.args				
Further arguments, ignored.		Named arguments to pass to storeDelayedObject.			
		Further arguments, ignored.			

# Value

x is saved to path and NULL is invisibly returned.

# Author(s)

Aaron Lun

# See Also

storeDelayedObject, for the methods to save each delayed operation.

#### saveSparseMatrix

# Examples

```
mat <- Matrix::rsparsematrix(100, 200, density=0.2)
rownames(mat) <- paste0("GENE_", seq_len(nrow(mat)))
dmat <- DelayedArray::DelayedArray(mat) * 1
dir <- tempfile()
saveObject(dmat, dir, delayed.preserve.ops=TRUE)
list.files(dir)</pre>
```

saveSparseMatrix Save a sparse matrix to disk

# Description

Save a sparse matrix to its on-disk representations.

# Usage

```
## S4 method for signature 'sparseMatrix'
saveObject(x, path, ...)
## S4 method for signature 'SVT_SparseMatrix'
saveObject(x, path, ...)
```

#### Arguments

х	A sparse matrix of some kind, typically from either the Matrix or SparseArray
	packages.
path	String containing the path to a directory in which to save x.
	Further arguments, currently ignored.

# Value

x is saved to path and NULL is invisibly returned.

# Author(s)

Aaron Lun

# See Also

readSparseMatrix, to read the directory contents back into the R session.

```
mat <- Matrix::rsparsematrix(100, 200, density=0.2)
rownames(mat) <- paste0("GENE_", seq_len(nrow(mat)))
dir <- tempfile()
saveObject(mat, dir)
list.files(dir)</pre>
```

storeDelayedObject Store/reload a DelayedArray

# Description

Store or reload the delayed operations or array-like seeds of a DelayedArray in an existing HDF5 file.

# Usage

```
storeDelayedObject(x, handle, name, ...)
reloadDelayedObject(handle, name, version = package_version("1.1"), ...)
reloadDelayedObjectFunctionRegistry(type = c("operation", "array"))
registerReloadDelayedObjectFunction(
  type = c("operation", "array"),
  subtype,
  fun,
  existing = c("old", "new", "error")
)
## S4 method for signature 'array'
storeDelayedObject(
  х,
  handle,
  name,
  version = package_version("1.1"),
  save.external.array = FALSE,
  . . .
)
## S4 method for signature 'ANY'
storeDelayedObject(
  х,
  handle,
  name,
  version = package_version("1.1"),
  external.save.args = list(),
  external.dedup.session = NULL,
  external.dedup.action = c("link", "copy", "symlink", "relsymlink"),
  . . .
)
altStoreDelayedObjectFunction(store)
altStoreDelayedObject(...)
altReloadDelayedObjectFunction(reload)
```

# storeDelayedObject

altReloadDelayedObject(...)

createExternalSeedDedupSession()

# Arguments

-	
х	Any of the delayed operation/array classes from DelayedArray.
handle	An <b>rhdf5</b> handle of a HDF5 file to save into (for storeDelayedObject) or load from (for reloadDelayedObject).
name	String containing the name of the group in file to save into (for storeDelayedObject) or load from (for reloadDelayedObject).
	For storeDelayedObject and reloadDelayedObject, additional arguments to be passed to specific methods.
	For altStoreDelayedObject and altReloadDelayedObject, arguments to be passed to the alternative functions.
version	Package version of the <b>chihaya</b> format to use when storing or reloading delayed objects. When reloading, the version should be retrieved from the attributes of the outermost group, typically by readDelayedArray.
type	String specifying the type of delayed object, i.e., operation or array. This corresponds to delayed_type type in the <b>chihaya</b> attributes.
subtype	String specifying the subtype of the delayed object, This corresponds to delayed_array or delayed_operation type (depending on type) in the <b>chihaya</b> attributes.
fun	Function to reload a delayed object. This should accept the same arguments as reloadDelayedObject and should return a delayed array (if type="array") or operation (otherwise). It may also be NULL to delete an existing entry in the registry.
existing	Logical scalar indicating the action to take if a function has already been regis- tered for type and subtype - keep the old or new function, or throw an error.
save.external.a	array
	Logical scalar indicating whether to save an array-like seed as an external seed, even if a dedicated storeDelayedObject method is available.
external.save.a	-
	Named list of further arguments to pass to altSaveObject when saving an ex- ternal seed.
external.dedup	.session Session object created by createExternalSeedDedupSession.
external.dedup	
	String specifying the deduplication method to use.
store	Function (typically a generic) to store delayed objects to file. This should accept the same arguments as storeDelayedObject.
reload	Function to reload delayed objects from file. This should accept the same arguments as reloadDelayedObject.

# Value

For storeDelayedObject and altStoreDelayedObject, the contents of x are saved to file, and NULL is invisibly returned.

For reloadDelayedObject and altReloadDelayedObject, a delayed operation or DelayedArray is returned.

For altStoreDelayedObjectFunction, the current store function is returned if store is missing. Otherwise, store is set as the current store function and the previous store function is returned.

For altReloadDelayedObjectFunction, the current reload function is returned if reload is missing. Otherwise, reload is set as the current reload function and the previous reload function is returned.

#### Customization

Developers can easily extend **alabaster.matrix** to new delayed objects by writing new methods for storeDelayedObject. Methods should save the contents of the delayed object to the HDF5 file in the **chihaya** format. Each new store method typically requires a corresponding reloading function to be registered via registerReloadDelayedObjectFunction, so that reloadDelayedObject knows how to reconstitute the object from file.

Application developers can customize the process of storing/reloading delayed objects by specifying alternative functions in altReloadDelayedObjectFunction and altStoreDelayedObjectFunction. For example, if we want to preserve all delayed objects except for DelayedSubset, we could replace storeDelayedObject with an altStoreDelayedObject that realizes any DelayedSubset instance into an ordinary matrix. This is analogous to the overrides for altReadObject and altSaveObject.

Extension developers (i.e., those who write new methods for storeDelayedObject or new functions for reloadDelayedObject) should generally use altStoreDelayedObject and altReloadDelayedObject in their method/funcion bodies. This ensures that any custom overrides specified by application developers are still respected in the extensions to **alabaster.matrix**.

#### **External seeds**

Whenever storeDelayedObject encounters a delayed operation or array-like seed for which it has no methods, the ANY method will save the delayed object as an "external seed". The array is saved via altSaveObject into a seeds directory next to the file associated with handle. A reference to this external location is then stored in the name group inside handle.

Users can force this behavior for all array-like seeds by passing save.external.array=TRUE in the ... arguments of storeDelayedObject. This instructs storeDelayedObject to save everything as external seeds, including those arrays for which it has methods. Doing so can be beneficial to enable deduplication, e.g., when two delayed arrays perform different operations on the same underlying seed. By saving the seeds externally, file management systems can identify the redundancy to save storage space.

Advanced users can explicitly deduplicate external seeds by setting save.external.array=TRUE and passing external.dedup.session= to storeDelayedObject. The external.dedup.session object is filled up with unique seeds as storeDelayedObject is called on various DelayedArrays. Whenever a duplicate seed is encountered, it is not saved again, but is instead linked or copied from the file path associated with the identical external seed. For example, a new session can be created when saving a SummarizedExperiment to deduplicate seeds across its assays.

The exact deduplication action can be specified by specifying the external.dedup.action= parameter. By default, storeDelayedObject attempts to create hard links, falling back to copies when a link cannot be created. Users can instead create copies, symbolic links to absolute paths, or even symbolic links to relative paths (e.g., to link to a "neighboring" assay of the same SummarizedExperiment).

When external seeds are encountered by reloadDelayedObject, they are loaded as ReloadedArrays (or some variant thereof) by altReadObject. Users can forcibly realize the reloaded seed into memory by passing custom.takane.reload=TRUE in ... for the reloadDelayedObject call. This is occasionally helpful for providing a more faithful roundtrip from file back into memory.

#### WrapperArraySeed

#### Author(s)

Aaron Lun

# See Also

saveObject,DelayedArray-method and readDelayedArray, where these methods are used. https://artifactdb.github.io/chihaya/, for the file format specification of delayed objects.

# Examples

```
library(DelayedArray)
X <- DelayedArray(matrix(runif(100), ncol=20))
Y <- cbind(X, DelayedArray::ConstantArray(value=50, c(5, 10)))
library(rhdf5)
temp <- tempfile()
dir.create(temp)
fpath <- file.path(temp, "foo.h5")
fhandle <- H5Fcreate(fpath)
storeDelayedObject(Y@seed, fhandle, "YAY")
rhdf5::h5ls(fhandle)
H5Fclose(fhandle)
fhandle <- H5Fopen(fpath, "H5F_ACC_RDONLY")
reloadDelayedObject(fhandle, "YAY")
H5Fclose(fhandle)
```

WrapperArraySeed DelayedArray wrapper seed

# Description

The WrapperArraySeed is, as the name suggests, a virtual class for a DelayedArray wrapper seed. This forwards most of the DelayedArray generic operations onto an internal seed class, typically a H5SparseMatrixSeed or HDF5ArraySeed objects from readSparseMatrix or readArray. Similarly, the WrapperArray is a virtual DelayedArray class that contains a WrapperArraySeed.

If an **alabaster** application operates on large arrays, developers may can consider defining concrete subclasses of the WrapperArraySeed (and WrapperArray). These subclasses can store application-specific provenance-tracking information that persist throughout the lifetime of the array. Such information is most useful for optimizing saveObject calls, which can instruct the application to link to the existing array rather than creating a new file. Check out the ReloadedArraySeed class for an example of this approach.

```
y <- Matrix::rsparsematrix(1000, 100, 0.01)
foo <- new("FooArraySeed", seed=y, foo_id="F00.0001")
dim(foo)
is_sparse(foo)
extract_array(foo, list(1:10, 1:10))
extract_sparse_array(foo, list(1:10, 1:10))</pre>
```

writeSparseMatrix Write a sparse matrix

# Description

Writes a sparse matrix to file in a compressed sparse format.

## Usage

```
writeSparseMatrix(
    x,
    file,
    name,
    chunk = 10000,
    column = TRUE,
    tenx = FALSE,
    guess.integer = TRUE
)
```

# Arguments

х	A sparse matrix of some sort. This includes sparse DelayedMatrix objects.
file	String containing a path to the HDF5 file. The file is created if it is not already present.
name	String containing the name of the group to store x.
chunk	Integer scalar specifying the chunk size for the indices and values.
column	Logical scalar indicating whether to store as compressed sparse column format.
tenx	Logical scalar indicating whether to use the 10X compressed sparse column format.
guess.integer	Logical scalar specifying whether to guess an appropriate integer type from x.

# Details

This writes a sparse matrix to file in various formats:

- column=TRUE and tenx=FALSE uses H5AD's csr\_matrix format.
- column=FALSE and tenx=FALSE uses H5AD's csc\_matrix format.
- tenx=TRUE uses 10X Genomics' HDF5 matrix format.

```
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```

#### writeSparseMatrix

For the first two formats, the apparent transposition is deliberate, because columns in R are interpreted as rows in H5AD. This allows us to retain consistency the interpretation of samples (columns in R, rows in H5AD) and features (vice versa). Constructors for classes like H5SparseMatrix will automatically transpose so no extra work is required.

If guess.integer=TRUE, we attempt to save x's values into the smallest type that will accommodate all of its values. If x only contains unsigned integers, we will attempt to save either 8-, 16- or 32-bit unsigned integers. If x contains signed integers, we will fall back to 32-bit signed integers. For all other values, we will fall back to double-precision floating point values.

We attempt to save x's indices to unsigned 16-bit integers if the relevant dimension of x is small enough. Otherwise we will save it as an unsigned 32-bit integer.

# Value

A NULL invisibly. The contents of x are written to name in file.

#### Author(s)

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```
library(Matrix)
x <- rsparsematrix(100, 20, 0.5)
tmp <- tempfile(fileext=".h5")
writeSparseMatrix(x, tmp, "csc_matrix")
writeSparseMatrix(x, tmp, "csr_matrix", column=FALSE)
writeSparseMatrix(x, tmp, "tenx_matrix", tenx = TRUE)
rhdf5::h5ls(tmp)
library(HDF5Array)
H5SparseMatrix(tmp, "csc_matrix")
H5SparseMatrix(tmp, "csr_matrix")</pre>
```

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