# Package 'coolR'

September 10, 2020
Title Convert Cooler HDF5 Contact Matrix To InteractionSet Objects
<b>Version</b> 0.0.0.9000
<b>Description</b> What the package does (one paragraph).
License `use_mit_license()`
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R topics documented:
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cool2IntSet Read .cool/.mcool sparse matrix into a GInteractions
<b>Description</b> Read .cool/.mcool sparse matrix into a GInteractions
Usage
<pre>cool2IntSet(files, res = NULL, cores = detectCores())</pre>
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2 read.cool

#### **Arguments**

files	A character vector of paths to HDF5 stored cool (uni-dimension) or mool (multi-dimension sparse matrix). If using a multi-dimesion, the resolution of one of the dimension need to be passed to res.
res	'NULL' if using a uni-dimensional cool file or the resolution of one of layer in the mcool file
cores	An integer for the number of parallel thread to convert the file

#### Value

An InteractionSet of all common pairs with the number of interaction for each pairs

getBins	Accessing the genomic bins in a cool/mcool file	

#### **Description**

Retrieves the chromosome bins forming from a HDF5 stored uni- or multi-dimentional contact matrix (cool or mcool file) into a seqinfo object

#### Usage

```
getBins(file, res)
```

# Arguments

Arguments	
file	Path to a HDF5 stored cool (uni-dimension) or mcool (multi-dimension sparse matrix). If using a multi-dimesion, the resolution of one of the dimension need to be passed to res
res	'NULL' if using a uni-dimensional cool file or the resolution of one of layer in the mcool file
read.cool	Read .cool/.mcool sparse matrix into a GInteractions

## Description

This function reads a compressed sparse row (CSR) storage scheme for a matrix created by the cooler application (https://github.com/mirnylab/cooler) stored in a HDF5 data storage. This utility can read the matrix or only specfic region of the genome either from one pair or a pair of region. If using a multi-dimension dataset (ie .mcool), the resolution needed need to be specified.

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# Usage

```
read.cool(
  file,
  res = NULL,
  chr1 = NULL,
  start1 = NULL,
  end1 = NULL,
  chr2 = NULL,
  start2 = NULL,
  end2 = NULL)
```

## Arguments

file	Path to a HDF5 stored cool (uni-dimension) or mcool (multi-dimension sparse matrix). If using a multi-dimesion, the resolution of one of the dimension need to be passed to res
res	'NULL' if using a uni-dimensional cool file or the resolution of one of layer in the mcool file
chr1	'NULL' or the chromosome name of the first pairs to return
start1	'NULL' or the start of a region within chr1 to return the pairs from. If chr1 is set and start1 is NULL this will be set to 1 by default
end1	'NULL' or the end of a region within chr1 to return the pairs from. If chr1 is set and end1 is NULL this will be set to the length of chr1 by default
chr2	'NULL' or the chromosome name of the first pairs to return
start2	'NULL' or the start of a region within chr2 to return the pairs from. If chr1 is set and start1 is NULL this will be set to 1 by default
end2	'NULL' or the end of a region within chr1 to return the pairs from. If chr2 is set and end1 is NULL this will be set to the length of chr2 by default

#### Value

A GInteraction object of the contact matrix dataset

seqinfo.cool	Accessing the sequence information in a cool/mcool file	

## Description

Convering the chromosomes information from a HDF5 stored uni- or multi-dimentional contact matrix (cool or mcool file) into a seqinfo object

## Usage

```
seqinfo.cool(file, res)
```

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# Arguments

file Path to a HDF5 stored cool (uni-dimension) or mcool (multi-dimension sparse

matrix). If using a multi-dimesion, the resolution of one of the dimension need

to be passed to res

res 'NULL' if using a uni-dimensional cool file or the resolution of one of layer in

the mcool file

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