

# Package ‘cmapR’

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**Description** The Connectivity Map (CMap) is a massive resource of perturbational gene expression profiles built by researchers at the Broad Institute and funded by the NIH Library of Integrated Network-Based Cellular Signatures (LINCS) program. Please visit <https://clue.io> for more information.  
The cmapR package implements methods to parse, manipulate, and write common CMap data objects, such as annotated matrices and collections of gene sets.

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---

**align\_matrices***Align the rows and columns of two (or more) matrices*

---

## Description

Align the rows and columns of two (or more) matrices

## Usage

```
align_matrices(m1, m2, ..., L = NULL, na.pad = TRUE, as.3D = TRUE)
```

## Arguments

m1	a matrix with unique row and column names
m2	a matrix with unique row and column names
...	additional matrices with unique row and column names
L	a list of matrix objects. If this is given, m1, m2, and ... are ignored
na.pad	boolean indicating whether to pad the combined matrix with NAs for rows/columns that are not shared by m1 and m2.
as.3D	boolean indicating whether to return the result as a 3D array. If FALSE, will return a list.

## Value

an object containing the aligned matrices. Will either be a list or a 3D array

## Examples

```
# construct some example matrices
m1 <- matrix(rnorm(20), nrow=4)
rownames(m1) <- letters[1:4]
colnames(m1) <- LETTERS[1:5]
m2 <- matrix(rnorm(20), nrow=5)
rownames(m2) <- letters[1:5]
colnames(m2) <- LETTERS[1:4]
m1
m2

# align them, padding with NA and returning a 3D array
align_matrices(m1, m2)

# align them, not padding and retuning a list
align_matrices(m1, m2, na.pad=FALSE, as.3D=FALSE)
```

`annotate.gct`      *Add annotations to a GCT object*

## Description

Given a GCT object and either a [data.frame](#) or a path to an annotation table, apply the annotations to the gct using the given keyfield.

## Usage

```
annotate.gct(...)

annotate_gct(g, annot, dim = "row", keyfield = "id")

## S4 method for signature 'GCT'
annotate_gct(g, annot, dim = "row", keyfield = "id")
```

## Arguments

...	arguments passed on to <code>annotate_gct</code>
<code>g</code>	a GCT object
<code>annot</code>	a <a href="#">data.frame</a> or path to text table of annotations
<code>dim</code>	either 'row' or 'column' indicating which dimension of <code>g</code> to annotate
<code>keyfield</code>	the character name of the column in <code>annot</code> that matches the row or column identifiers in <code>g</code>

## Value

a GCT object with annotations applied to the specified dimension

## See Also

Other GCT utilities: [melt.gct\(\)](#), [merge.gct\(\)](#), [rank.gct\(\)](#), [subset.gct\(\)](#)

## Examples

```
gct_path <- system.file("extdata", "modzs_n25x50.gctx", package="cmapR")
# read the GCT file, getting the matrix only
g <- parse_gctx(gct_path, matrix_only=TRUE)
# separately, read the column annotations and then apply them using
# annotate_gct
cdesc <- read_gctx_meta(gct_path, dim="col")
g <- annotate_gct(g, cdesc, dim="col", keyfield="id")
```

---

cdesc\_char

*An example table of metadata, as would be parsed from or parse.gctx.  
Initially all the columns are of type character.*

---

## Description

An example table of metadata, as would be parsed from or parse.gctx. Initially all the columns are of type character.

## Usage

cdesc\_char

## Format

An object of class `data.frame` with 368 rows and 8 columns.

---

check\_colnames

*Check whether test\_names are columns in the `data.frame` df*

---

## Description

Check whether `test_names` are columns in the `data.frame` `df`

## Usage

`check_colnames(test_names, df, throw_error = TRUE)`

## Arguments

<code>test_names</code>	a vector of column names to test
<code>df</code>	the <code>data.frame</code> to test against
<code>throw_error</code>	boolean indicating whether to throw an error if any <code>test_names</code> are not found in <code>df</code>

## Value

boolean indicating whether or not all `test_names` are columns of `df`

## Examples

```
check_colnames(c("pert_id", "pert_iname"), cdesc_char) # TRUE  
check_colnames(c("pert_id", "foobar"),  
              cdesc_char, throw_error=FALSE) # FALSE, suppress error
```

<code>check_dups</code>	<i>Check for duplicates in a vector</i>
-------------------------	---

### Description

Check for duplicates in a vector

### Usage

```
check_dups(x, name = "")
```

### Arguments

<code>x</code>	the vector
<code>name</code>	the name of the object to print in an error message if duplicates are found

### Value

silently returns NULL

### Examples

```
# this will throw an error, let's catch it
tryCatch(
  check_dups(c("a", "b", "c", "a", "d")),
  error=function(e) print(e)
)
```

<code>distil</code>	<i>Collapse the rows or columns of a matrix using weighted averaging</i>
---------------------	--

### Description

This is equivalent to the 'modz' procedure used in collapsing replicates in traditional L1000 data processing. The weight for each replicate is computed as its normalized average correlation to the other replicates in the set.

### Usage

```
distil(m, dimension = "col", method = "spearman")
```

### Arguments

<code>m</code>	a numeric matrix where the rows or columns are assumed to be replicates
<code>dimension</code>	the dimension to collapse. either 'row' or 'col'
<code>method</code>	the correlation method to use

**Value**

a list with the following elements

**values** a vector of the collapsed values

**correlations** a vector of the pairwise correlations

**weights** a vector of the computed weights

**Examples**

```
m <- matrix(rnorm(30), ncol=3)
distil(m)
```

---

ds

*An example of a GCT object with row and column metadata and gene expression values in the matrix.*

---

**Description**

An example of a GCT object with row and column metadata and gene expression values in the matrix.

**Usage**

ds

**Format**

An object of class GCT of length 1.

---

extract.gct

*Extract elements from a GCT matrix*

---

**Description**

extract the elements from a GCT object where the values of `row_field` and `col_field` are the same. A concrete example is if `g` represents a matrix of signatures of genetic perturbations, and you want to extract all the values of the targeted genes.

**Usage**

```
extract.gct(...)

extract_gct(
  g,
  row_field,
  col_field,
  rdesc = NULL,
  cdesc = NULL,
  row_keyfield = "id",
  col_keyfield = "id"
)
```

**Arguments**

...	arguments passed on to extract_gct
g	the GCT object
row_field	the column name in rdesc to search on
col_field	the column name in cdesc to search on
rdesc	a data.frame of row annotations
cdesc	a data.frame of column annotations
row_keyfield	the column name of rdesc to use for annotating the rows of g
col_keyfield	the column name of cdesc to use for annotating the rows of g

**Value**

a list of the following elements

**mask** a logical matrix of the same dimensions as ds@mat indicating which matrix elements have been extracted

**idx** an array index into ds@mat representing which elements have been extracted

**vals** a vector of the extracted values

**Examples**

```
# get the values for all targeted genes from a
# dataset of knockdown experiments
res <- extract_gct(kd_gct, row_field="pr_gene_symbol",
  col_field="pert_mfc_desc")
str(res)
stats::quantile(res$vals)
```

---

GCT	<i>Initialize an object of class GCT</i>
-----	--

---

## Description

Initialize an object of class GCT

## Usage

```
GCT(  
  mat = NULL,  
  rdesc = NULL,  
  cdesc = NULL,  
  src = NULL,  
  rid = NULL,  
  cid = NULL,  
  matrix_only = FALSE  
)
```

## Arguments

mat	a matrix
rdesc	a <code>data.frame</code> of row metadata
cdesc	a <code>data.frame</code> of column metadata
src	path to a GCT file to read
rid	vector of character identifiers for rows
cid	vector of character identifiers for columns
matrix_only	logical indicating whether to read just the matrix data from <code>src</code>

## Details

If `mat` is provided, `rid` and `cid` are treated as the row and column identifiers for the matrix and are assigned to the `rid` and `cid` slots of the GCT object.

If `mat` is not provided but `src` is provided, `rid` and `cid` are treated as filters. Data will be read from the file path provided to `src` and will then be restricted to the character ids or integer indices provided to `rid` and `cid`. In a similar manner, `matrix_only` controls whether the row and column metadata are also read from the `src` file path.

## Value

a GCT object

## See Also

Other GCTX parsing functions: [append.dim\(\)](#), [fix.datatypes\(\)](#), [parse.gctx\(\)](#), [process\\_ids\(\)](#), [read.gctx.ids\(\)](#), [read.gctx.meta\(\)](#), [write.gctx.meta\(\)](#), [write.gctx\(\)](#), [write.gct\(\)](#)

## Examples

```
# an empty object
(g <- GCT())
# with a matrix
# note we must specify row and column ids
(g <- GCT(mat=matrix(rnorm(100), nrow=10),
           rid=letters[1:10], cid=letters[1:10]))
# from file
gct_file <- system.file("extdata", "modzs_n25x50.gctx", package="cmapR")
(g <- GCT(src=gct_file))
```

**GCT-class**

*An S4 class to represent a GCT object*

## Description

The GCT class serves to represent annotated matrices. The `mat` slot contains said data and the `rdesc` and `cdesc` slots contain data frames with annotations about the rows and columns, respectively

## Slots

- `mat` a numeric matrix
- `rid` a character vector of row ids
- `cid` a character vector of column ids
- `rdesc` a `data.frame` of row descriptors
- `cdesc` a `data.frame` of column descriptors
- `src` a character indicating the source (usually file path) of the data

## See Also

[parse\\_gctx](#), [write\\_gctx](#), [read\\_gctx\\_meta](#), [read\\_gctx\\_ids](#)  
visit <http://clue.io/help> for more information on the GCT format

**gene\_set**

*An example collection of gene sets as used in the Lamb 2006 CMap paper.*

## Description

An example collection of gene sets as used in the Lamb 2006 CMap paper.

## Usage

`gene_set`

**Format**

An object of class `list` of length 8.

**Source**

Lamb et al 2006 doi:10.1126/science.1132939

---

ids	<i>Extract the or set row or column ids of a GCT object</i>
-----	---

---

**Description**

Extract the or set row or column ids of a GCT object

**Usage**

```
ids(g, dimension = "row")

## S4 method for signature 'GCT'
ids(g, dimension = "row")

ids(g, dimension = "row") <- value

## S4 replacement method for signature 'GCT'
ids(g, dimension = "row") <- value
```

**Arguments**

g	the GCT object
dimension	the dimension to extract/update ['row' or 'column']
value	a character vector

**Value**

a vector of row ids

**See Also**

Other GCT accessor methods: `mat()`, `meta()`

## Examples

```
# extract rids
rids <- ids(ds)
# extract column ids
cids <- ids(ds, "column")
# set rids
ids(ds) <- as.character(1:length(rids))
# set cids
ids(ds, "column") <- as.character(1:length(cids))
```

**is.wholenumber**      *Check if x is a whole number*

## Description

Check if x is a whole number

## Usage

```
is.wholenumber(x, tol = .Machine$double.eps^0.5)
```

## Arguments

x	number to test
tol	the allowed tolerance

## Value

boolean indicating whether x is tol away from a whole number value

## Examples

```
is.wholenumber(1)
is.wholenumber(0.5)
```

**kd\_gct**      *An example GCT object of knockdown experiments targeting a subset of landmark genes.*

## Description

An example GCT object of knockdown experiments targeting a subset of landmark genes.

## Usage

```
kd_gct
```

## Format

An object of class GCT of length 1.

---

lxb2mat

*Read an LXB file and return a matrix*

---

## Description

Read an LXB file and return a matrix

## Usage

```
lxb2mat(lxb_path, columns = c("RID", "RP1"), newnames = c("barcode_id", "FI"))
```

## Arguments

lxb_path	the path to the lxb file
columns	which columns in the lxb file to retain
newnames	what to name these columns in the returned matrix

## Value

a matrix

## See Also

Other CMap parsing functions: [parse.gmt\(\)](#), [parse.gmx\(\)](#), [parse.grp\(\)](#), [write\\_gmt\(\)](#), [write\\_grp\(\)](#)

## Examples

```
lxb_path <- system.file("extdata", "example.lxb", package="cmapR")
lxb_data <- lxb2mat(lxb_path)
str(lxb_data)
```

<code>mat</code>	<i>Extract or set the matrix of GCT object</i>
------------------	--

## Description

Extract or set the matrix of GCT object

## Usage

```
mat(g)

## S4 method for signature 'GCT'
mat(g)

mat(g) <- value

## S4 replacement method for signature 'GCT'
mat(g) <- value
```

## Arguments

<code>g</code>	the GCT object
<code>value</code>	a numeric matrix

## Value

a matrix

## See Also

Other GCT accessor methods: [ids\(\)](#), [meta\(\)](#)

## Examples

```
# get the matrix
m <- mat(ds)
# set the matrix
mat(ds) <- matrix(0, nrow=nrow(m), ncol=ncol(m))
```

---

<code>melt.gct</code>	<i>Transform a GCT object in to a long form <a href="#">data.table</a> (aka 'melt')</i>
-----------------------	---

---

## Description

Utilizes the `melt.data.table` function to transform the matrix into long form. Optionally can include the row and column annotations in the transformed `data.table`.

## Usage

```

melt.gct(...)

melt_gct(
  g,
  suffixes = NULL,
  remove_symmetries = FALSE,
  keep_rdesc = TRUE,
  keep_cdesc = TRUE,
  ...
)

## S4 method for signature 'GCT'
melt_gct(
  g,
  suffixes = NULL,
  remove_symmetries = FALSE,
  keep_rdesc = TRUE,
  keep_cdesc = TRUE,
  ...
)

```

## Arguments

...	further arguments passed along to <code>data.table::merge</code>
<code>g</code>	the GCT object
<code>suffixes</code>	the character suffixes to be applied if there are collisions between the names of the row and column descriptors
<code>remove_symmetries</code>	boolean indicating whether to remove the lower triangle of the matrix (only applies if <code>g@mat</code> is symmetric)
<code>keep_rdesc</code>	boolean indicating whether to keep the row descriptors in the final result
<code>keep_cdesc</code>	boolean indicating whether to keep the column descriptors in the final result

## Value

a `data.table` object with the row and column ids and the matrix values and (optionally) the row and column descriptors

**See Also**

Other GCT utilities: [annotate.gct\(\)](#), [merge.gct\(\)](#), [rank.gct\(\)](#), [subset.gct\(\)](#)

**Examples**

```
# simple melt, keeping both row and column meta
head(melt_gct(ds))

# update row/column suffixes to indicate rows are genes, columns experiments
head(melt_gct(ds, suffixes = c("_gene", "_experiment")))

# ignore row/column meta
head(melt_gct(ds, keep_rdesc = FALSE, keep_cdesc = FALSE))
```

**merge.gct**

*Merge two GCT objects together*

**Description**

Merge two GCT objects together

**Usage**

```
## S3 method for class 'gct'
merge(...)

merge_gct(g1, g2, dim = "row", matrix_only = FALSE)

## S4 method for signature 'GCT,GCT'
merge_gct(g1, g2, dim = "row", matrix_only = FALSE)
```

**Arguments**

...	arguments passed on to merge_gct
g1	the first GCT object
g2	the second GCT object
dim	the dimension on which to merge (row or column)
matrix_only	boolean indicating whether to keep only the data matrices from g1 and g2 and ignore their row and column meta data

**Value**

a GCT object

**See Also**

Other GCT utilities: [annotate.gct\(\)](#), [melt.gct\(\)](#), [rank.gct\(\)](#), [subset.gct\(\)](#)

## Examples

```
# take the first 10 and last 10 rows of an object
# and merge them back together
(a <- subset_gct(ds, rid=1:10))
(b <- subset_gct(ds, rid=969:978))
(merged <- merge_gct(a, b, dim="row"))
```

---

meta	<i>Extract the or set metadata of a GCT object</i>
------	--

---

## Description

Extract the or set metadata of a GCT object

## Usage

```
meta(g, dimension = "row")

## S4 method for signature 'GCT'
meta(g, dimension = "row")

meta(g, dimension = "row") <- value

## S4 replacement method for signature 'GCT'
meta(g, dimension = "row") <- value
```

## Arguments

g	the GCT object
dimension	the dimension to extract/update ['row' or 'column']
value	a data.frame

## Value

a data.frame

## See Also

Other GCT accessor methods: [ids\(\)](#), [mat\(\)](#)

## Examples

```
# extract rdesc
rdesc <- meta(ds)
# extract cdesc
cdesc <- meta(ds, dim="column")
# set rdesc
meta(ds) <- data.frame(x=sample(letters, nrow(rdesc), replace=TRUE))
# set cdesc
meta(ds, dim="column") <- data.frame(x=sample(letters, nrow(cdesc),
replace=TRUE))
```

**na\_pad\_matrix**

*Pad a matrix with additional rows/columns of NA values*

## Description

Pad a matrix with additional rows/columns of NA values

## Usage

```
na_pad_matrix(m, row_universe = NULL, col_universe = NULL)
```

## Arguments

- m a matrix with unique row and column names
- row\_universe a vector with the universe of possible row names
- col\_universe a vector with the universe of possible column names

## Value

a matrix

## Examples

```
m <- matrix(rnorm(10), nrow=2)
rownames(m) <- c("A", "B")
colnames(m) <- letters[1:5]
na_pad_matrix(m, row_universe=LETTERS, col_universe=letters)
```

---

`parse.gctx`

*Parse a GCTX file into the workspace as a GCT object*

---

## Description

Parse a GCTX file into the workspace as a GCT object

## Usage

```
parse.gctx(...)
```

```
parse_gctx(fname, rid = NULL, cid = NULL, matrix_only = FALSE)
```

## Arguments

...	arguments passed on to <code>parse_gctx</code>
fname	path to the GCTX file on disk
rid	either a vector of character or integer row indices or a path to a grp file containing character row indices. Only these indicies will be parsed from the file.
cid	either a vector of character or integer column indices or a path to a grp file containing character column indices. Only these indicies will be parsed from the file.
matrix_only	boolean indicating whether to parse only the matrix (ignoring row and column annotations)

## Details

`parse_gctx` also supports parsing of plain text GCT files, so this function can be used as a general GCT parser.

## Value

a GCT object

## See Also

Other GCTX parsing functions: [GCT](#), [append.dim\(\)](#), [fix.datatypes\(\)](#), [process\\_ids\(\)](#), [read.gctx.ids\(\)](#), [read.gctx.meta\(\)](#), [write.gctx.meta\(\)](#), [write.gctx\(\)](#), [write.gct\(\)](#)

## Examples

```
gct_file <- system.file("extdata", "modzs_n25x50.gctx", package="cmapR")
(ds <- parse_gctx(gct_file))

# matrix only
(ds <- parse_gctx(gct_file, matrix_only=TRUE))
```

---

```
# only the first 10 rows and columns
(ds <- parse_gctx(gct_file, rid=1:10, cid=1:10))
```

---

**parse.gmt***Read a GMT file and return a list***Description**

Read a GMT file and return a list

**Usage**

```
parse.gmt(. . .)
parse_gmt(fname)
```

**Arguments**

. . .	arguments passed on to parse_gmt
fname	the file path to be parsed

**Details**

`parse_gmt` returns a nested list object. The top level contains one list per row in `fname`. Each of these is itself a list with the following fields: - `head`: the name of the data (row in `fname`) - `desc`: description of the corresponding data - `len`: the number of data items - `entry`: a vector of the data items

**Value**

a list of the contents of `fname`. See details.

**See Also**

Visit <http://clue.io/help> for details on the GMT file format

Other CMap parsing functions: `lxb2mat()`, `parse.gmx()`, `parse.grp()`, `write_gmt()`, `write_grp()`

**Examples**

```
gmt_path <- system.file("extdata", "query_up.gmt", package="cmapR")
gmt <- parse_gmt(gmt_path)
str(gmt)
```

---

parse.gmx

*Read a GMX file and return a list*

---

## Description

Read a GMX file and return a list

## Usage

```
parse.gmx(...)
```

```
parse_gmx(fname)
```

## Arguments

...	arguments passed on to parse_gmx
fname	the file path to be parsed

## Details

parse\_gmx returns a nested list object. The top level contains one list per column in fname. Each of these is itself a list with the following fields:

- head: the name of the data (column in fname)
- desc: description of the corresponding data
- len: the number of data items
- entry: a vector of the data items

## Value

a list of the contents of fname. See details.

## See Also

Visit <http://clue.io/help> for details on the GMX file format

Other CMap parsing functions: `lxb2mat()`, `parse.gmt()`, `parse.grp()`, `write_gmt()`, `write_grp()`

## Examples

```
gmx_path <- system.file("extdata", "lmb_probes.gmx", package="cmapR")
gmx <- parse_gmx(gmx_path)
str(gmx)
```

`parse.grp`*Read a GRP file and return a vector of its contents***Description**

Read a GRP file and return a vector of its contents

**Usage**

```
parse.grp(...)

parse_grp(fname)
```

**Arguments**

<code>...</code>	arguments passed on to <code>parse_grp</code>
<code>fname</code>	the file path to be parsed

**Value**

a vector of the contents of `fname`

**See Also**

Visit <http://clue.io/help> for details on the GRP file format

Other CMap parsing functions: `lxb2mat()`, `parse.gmt()`, `parse.gmx()`, `write_gmt()`, `write_grp()`

**Examples**

```
grp_path <- system.file("extdata", "lm_epsilon_n978.grp", package="cmapR")
values <- parse_grp(grp_path)
str(values)
```

`rank.gct`*Convert a GCT object's matrix to ranks***Description**

Convert a GCT object's matrix to ranks

**Usage**

```
rank.gct(...)

rank_gct(g, dim = "col", decreasing = TRUE)

## S4 method for signature 'GCT'
rank_gct(g, dim = "col", decreasing = TRUE)
```

**Arguments**

...	arguments passed on to <code>rank_gct</code>
<code>g</code>	the GCT object to rank
<code>dim</code>	the dimension along which to rank (row or column)
<code>decreasing</code>	boolean indicating whether higher values should get lower ranks

**Value**

a modified version of `g`, with the values in the matrix converted to ranks

**See Also**

Other GCT utilities: [annotate.gct\(\)](#), [melt.gct\(\)](#), [merge.gct\(\)](#), [subset.gct\(\)](#)

**Examples**

```
(ranked <- rank_gct(ds, dim="column"))
# scatter rank vs. score for a few columns
m <- mat(ds)
m_ranked <- mat(ranked)
plot(m[, 1:3], m_ranked[, 1:3],
     xlab="score", ylab="rank")
```

`read.gctx.ids` *Read GCTX row or column ids*

**Description**

Read GCTX row or column ids

**Usage**

```
read.gctx.ids(...)

read_gctx_ids(gctx_path, dim = "row")
```

**Arguments**

...	arguments passed on to <code>read_gctx_ids</code>
<code>gctx_path</code>	path to the GCTX file
<code>dim</code>	which ids to read (row or column)

**Value**

a character vector of row or column ids from the provided file

**See Also**

Other GCTX parsing functions: [GCT](#), [append.dim\(\)](#), [fix.datatypes\(\)](#), [parse.gctx\(\)](#), [process\\_ids\(\)](#), [read.gctx.meta\(\)](#), [write.gctx.meta\(\)](#), [write.gctx\(\)](#), [write.gct\(\)](#)

**Examples**

```
gct_file <- system.file("extdata", "modzs_n25x50.gctx", package="cmapR")
# row ids
rid <- read_gctx_ids(gct_file)
head(rid)
# column ids
cid <- read_gctx_ids(gct_file, dim="column")
head(cid)
```

<code>read.gctx.meta</code>	<i>Parse row or column metadata from GCTX files</i>
-----------------------------	---

**Description**

Parse row or column metadata from GCTX files

**Usage**

```
read.gctx.meta(...)

read_gctx_meta(gctx_path, dim = "row", ids = NULL)
```

**Arguments**

...	arguments passed on to <code>read_gctx_meta</code>
<code>gctx_path</code>	the path to the GCTX file
<code>dim</code>	which metadata to read (row or column)
<code>ids</code>	a character vector of a subset of row/column ids for which to read the metadata

**Value**

a `data.frame` of metadata

**See Also**

Other GCTX parsing functions: [GCT](#), [append.dim\(\)](#), [fix.datatypes\(\)](#), [parse.gctx\(\)](#), [process\\_ids\(\)](#), [read.gctx.ids\(\)](#), [write.gctx.meta\(\)](#), [write.gctx\(\)](#), [write.gct\(\)](#)

## Examples

```

gct_file <- system.file("extdata", "modzs_n25x50.gctx", package="cmapR")
# row meta
row_meta <- read_gctx_meta(gct_file)
str(row_meta)
# column meta
col_meta <- read_gctx_meta(gct_file, dim="column")
str(col_meta)
# now for only the first 10 ids
col_meta_first10 <- read_gctx_meta(gct_file, dim="column",
ids=col_meta$id[1:10])
str(col_meta_first10)

```

**robust\_zscore**      *Compute robust z-scores*

## Description

robust zscore implementation takes in a 1D vector, returns 1D vector after computing robust zscores  
 $rZ = (x - \text{med}(x)) / \text{mad}(x)$

## Usage

```
robust_zscore(x, min_mad = 1e-06, ...)
```

## Arguments

<code>x</code>	numeric vector to z-score
<code>min_mad</code>	the minimum allowed MAD, useful for avoiding division by very small numbers
<code>...</code>	further options to median, max functions

## Value

transformed version of `x`

## Examples

```

(x <- rnorm(25))
(robust_zscore(x))

# with min_mad
(robust_zscore(x, min_mad=1e-4))

```

**subset.gct***Subset a gct object using the provided row and column ids***Description**

Subset a gct object using the provided row and column ids

**Usage**

```
## S3 method for class 'gct'
subset(...)

subset_gct(g, rid = NULL, cid = NULL)

## S4 method for signature 'GCT'
subset_gct(g, rid = NULL, cid = NULL)
```

**Arguments**

...	arguments passed on to subset_gct
g	a gct object
rid	a vector of character ids or integer indices for ROWS
cid	a vector of character ids or integer indices for COLUMNS

**Value**

a GCT object

**See Also**

Other GCT utilities: [annotate.gct\(\)](#), [melt.gct\(\)](#), [merge.gct\(\)](#), [rank.gct\(\)](#)

**Examples**

```
# first 10 rows and columns by index
(a <- subset_gct(ds, rid=1:10, cid=1:10))

# first 10 rows and columns using character ids
# use \code{ids} to extract the ids
rid <- ids(ds)
cid <- ids(ds, dimension="col")
(b <- subset_gct(ds, rid=rid[1:10], cid=cid[1:10]))

identical(a, b) # TRUE
```

---

threshold	<i>Threshold a numeric vector</i>
-----------	-----------------------------------

---

**Description**

Threshold a numeric vector

**Usage**

```
threshold(x, minval, maxval)
```

**Arguments**

x	the vector
minval	minimum allowed value
maxval	maximum allowed value

**Value**

a thresholded version of x

**Examples**

```
x <- rnorm(20)
threshold(x, -0.1, -0.1)
```

---

---

transpose.gct	<i>Transpose a GCT object</i>
---------------	-------------------------------

---

**Description**

Transpose a GCT object

**Usage**

```
transpose.gct(...)
transpose_gct(g)

## S4 method for signature 'GCT'
transpose_gct(g)
```

**Arguments**

...	arguments passed on to transpose_gct
g	the GCT object

**Value**

a modified version of the input GCT object where the matrix has been transposed and the row and column ids and annotations have been swapped.

**Examples**

```
transpose_gct(ds)
```

---

update.gctx	<i>Update the matrix of an existing GCTX file</i>
-------------	---

---

**Description**

Update the matrix of an existing GCTX file

**Usage**

```
## S3 method for class 'gctx'
update(...)

update_gctx(x, ofile, rid = NULL, cid = NULL)
```

**Arguments**

...	arguments passed on to update_gctx
x	an array of data
ofile	the filename of the GCTX to update
rid	integer indices or character ids of the rows to update
cid	integer indices or character ids of the columns to update

**Details**

Overwrite the rows and columns of ofile as indicated by rid and cid respectively. rid and cid can either be integer indices or character ids corresponding to the row and column ids in ofile.

**Value**

silently returns NULL

## Examples

```
## Not run:
m <- matrix(rnorm(20), nrow=10)
# update by integer indices
update_gctx(m, ofile="my.gctx", rid=1:10, cid=1:2)
# update by character ids
row_ids <- letters[1:10]
col_ids <- LETTERS[1:2]
update_gctx(m, ofile="my.gctx", rid=row_ids, cid=col_ids)

## End(Not run)
```

**write.gct**

*Write a GCT object to disk in GCT format*

## Description

Write a GCT object to disk in GCT format

## Usage

```
write.gct(...)

write_gct(ds, ofile, precision = 4, appenddim = TRUE, ver = 3)
```

## Arguments

...	arguments passed on to write_gct
ds	the GCT object
ofile	the desired output filename
precision	the numeric precision at which to save the matrix. See details.
appenddim	boolean indicating whether to append matrix dimensions to filename
ver	the GCT version to write. See details.

## Details

Since GCT is text format, the higher precision you choose, the larger the file size. ver is assumed to be 3, aka GCT version 1.3, which supports embedded row and column metadata in the GCT file. Any other value passed to ver will result in a GCT version 1.2 file which contains only the matrix data and no annotations.

## Value

silently returns NULL

## See Also

Other GCTX parsing functions: [GCT](#), [append.dim\(\)](#), [fix.datatypes\(\)](#), [parse.gctx\(\)](#), [process\\_ids\(\)](#), [read.gctx.ids\(\)](#), [read.gctx.meta\(\)](#), [write.gctx.meta\(\)](#), [write.gctx\(\)](#)

## Examples

```
# note this will create a GCT file in your current directory
write_gct(ds, "dataset", precision=2)
```

### `write.gctx`

*Write a GCT object to disk in GCTX format*

## Description

Write a GCT object to disk in GCTX format

## Usage

```
write.gctx(...)

write_gctx(
    ds,
    ofile,
    appenddim = TRUE,
    compression_level = 0,
    matrix_only = FALSE,
    max_chunk_kb = 1024
)
```

## Arguments

...	arguments passed on to <code>write_gctx</code>
ds	a GCT object
ofile	the desired file path for writing
appenddim	boolean indicating whether the resulting filename will have dimensions appended (e.g. my_file_n384x978.gctx)
compression_level	integer between 1-9 indicating how much to compress data before writing. Higher values result in smaller files but slower read times.
matrix_only	boolean indicating whether to write only the matrix data (and skip row, column annotations)
max_chunk_kb	for chunking, the maximum number of KB a given chunk will occupy

**Value**

silently returns NULL

**See Also**

Other GCTX parsing functions: [GCT](#), [append.dim\(\)](#), [fix.datatypes\(\)](#), [parse.gctx\(\)](#), [process\\_ids\(\)](#), [read.gctx.ids\(\)](#), [read.gctx.meta\(\)](#), [write.gctx.meta\(\)](#), [write.gct\(\)](#)

**Examples**

```
# note this will create a GCT file in your current directory
write_gctx(ds, "dataset")
```

---

**write.tbl***Write a data.frame to a tab-delimited text file*

---

**Description**

Write a `data.frame` to a tab-delimited text file

**Usage**

```
write.tbl(...)
write_tbl(tbl, ofile, ...)
```

**Arguments**

...	additional arguments passed on to <code>write.table</code>
tbl	the <code>data.frame</code> to be written
ofile	the desired file name

**Details**

This method simply calls `write.table` with some preset arguments that generate a unquoted, tab-delimited file without row names.

**Value**

silently returns NULL

**See Also**

[write.table](#)

## Examples

```
## Not run:
write_tbl(cdesc_char, "col_meta.txt")

## End(Not run)
```

`write_gmt`

*Write a nested list to a GMT file*

## Description

Write a nested list to a GMT file

## Usage

```
write_gmt(lst, fname)
```

## Arguments

<code>lst</code>	the nested list to write. See details.
<code>fname</code>	the desired file name

## Details

`lst` needs to be a nested list where each sub-list is itself a list with the following fields: - `head`: the name of the data - `desc`: description of the corresponding data - `len`: the number of data items - `entry`: a vector of the data items

## Value

silently returns NULL

## See Also

Visit <http://clue.io/help> for details on the GMT file format

Other CMap parsing functions: `lxb2mat()`, `parse.gmt()`, `parse.gmx()`, `parse.grp()`, `write_grp()`

## Examples

```
## Not run:
write_gmt(gene_set, "gene_set.gmt")

## End(Not run)
```

---

`write_grp`*Write a vector to a GRP file*

---

## Description

Write a vector to a GRP file

## Usage

```
write_grp(vals, fname)
```

## Arguments

vals	the vector of values to be written
fname	the desired file name

## Value

silently returns NULL

## See Also

Visit <http://clue.io/help> for details on the GRP file format

Other CMap parsing functions: `lxb2mat()`, `parse.gmt()`, `parse.gmx()`, `parse.grp()`, `write_gmt()`

## Examples

```
## Not run:  
write_grp(letters, "letter.grp")  
  
## End(Not run)
```

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